

## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification <sup>6</sup>:
C12N 15/87, 5/10, A61K 9/48, 38/16, 38/33

(11) International Publication Number:

WO 96/40959

. **A1** 

(43) International Publication Date:

19 December 1996 (19.12.96)

(21) International Application Number:

PCT/US96/09629

(22) International Filing Date:

7 June 1996 (07.06.96)

(30) Priority Data:

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08/481,917

7 June 1995 (07.06.95)

US

(60) Parent Application or Grant

(63) Related by Continuation US

08/481,917 (CIP)

Filed on

7 June 1995 (07.06.95)

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(81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

#### Published

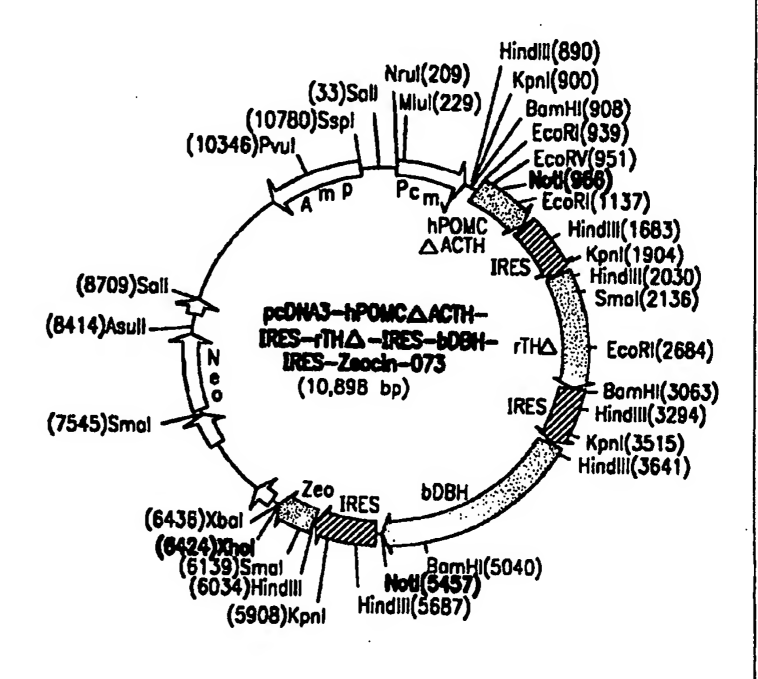
With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: CELL LINE PRODUCING ANALGESIC COMPOUNDS FOR TREATING PAIN

#### (57) Abstract

A genetically engineered cell line that produces at least one catecholamine, at least one endorphin, and at least one enkephalin, for the treatment of pain. The cells may be provided directly to a patient in need thereof, or encapsulated to form a bioartificial organ.



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Cell line producing analgesic compounds for treating pain

# Field of the Invention

The present invention relates to a cell line useful for the treatment of pain. More particularly, the cell line of this invention has been genetically engineered to produce at least one analgesic compound from each of the groups consisting of endorphins, enkephalins, and catecholamines.

### 10 Background of the Invention

Pain is a common symptom of disease. The superficial dorsal horn of the spinal cord, where primary afferent fibers carrying nociceptive information terminate, contains enkephalinergic interneurons and high densities of opiate receptors. In addition, there is a dense concentration of noradrenergic fibers in the superficial laminae of the spinal cord.

Acute pain arises in response to acute
noxious stimuli. Chronic pain is predominantly due to
neuropathies of central or peripheral origin. This

neuropathic pain is the result of aberrant somatosensory processing that can result in increased sensitivity to a painful stimulus (hyperalgesia) and pain associated with a stimulus that does not usually provoke pain (allodynia).

Intrathecal injection of morphine into the spinal subarachnoid space produces potent analgesia. Similarly, intrathecal administration of norepinephrine or noradrenergic agonists also produces analgesia.

10 See, e.g., Sagen et al., Proc. Natl. Acad. Sci. USA,
83, pp. 7522-26 (1986).

Co-administration of subeffective doses of opiates, such as enkephalins, and catecholamines, such as norepinephrine, may synergize to produce analgesia.

15 Ibid. Chromaffin cells in the adrenal medulla produce and release several neuroactive substances including norepinephrine, epinephrine, met-enkephalin, leuenkephalin, neuropeptide Y, vasoactive intestinal polypeptide, somatostatin, neurotensin, cholecystokinin and calcitonin gene-related peptide. See, e.g., Sagen et al., Proc. Natl. Acad. Sci. USA, 83, pp. 7522-26 (1986); Sagen et al., Jour. Neurochem., 56, pp. 623-27 (1991).

Because chromaffin cells produce both opioid

25 peptides and catecholamines, one approach to reduction
of nociceptive response or pain sensitivity has
investigated transplanting adrenal medullary tissue, as
well as isolated adrenal chromaffin cells, directly
into CNS pain modulatory regions, in attempts to

30 provide analgesia. See, e.g., Sagen et al., Brain
Research, 384, pp. 189-94 (1986); Vaguero et al.,
Neuroreport, 2, pp. 149-51 (1991); Ginzberg and

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Seltzer, Brain Research, 523, pp. 147-50 (1990); Sagen et al., Pain, 42. pp. 69-79 (1990).

Attempts to produce analgesic have been made using both allogeneic and xenogeneic chromaffin tissue or cells transplants. Allograft tissue is in limited supply, and is not readily available, particularly for in human pain treatment programs. In addition, allogeneic human tissue carries the risk of pathogenic contamination. See e.g., Hama and Sagen, Brain

Research, 651, pp. 183-93 (1994).

Xenogeneic donors may provide large quantities of material that can be readily obtained. For this reason, bovine adrenal tissue has been used. See, e.g., Hama and Sagen, Brain Research, 651, pp. 183-93 (1994).

However, potentially serious host
consequences, as well as ultimate graft rejection, are
inherent problems in transplantation between disparate
species. Complete graft rejection of whole or
dissociated tissue may occur even in the CNS, normally
thought to be immunologically privileged, due to
presence of highly antigenic cells in the xenografts,
particularly endothelial cells. In addition, the donor
tissue must be carefully screened to avoid introduction
of viral contaminants, or other pathogens, to the host.
To overcome graft rejection, immunosuppression is
required typically using cyclosporine A.

Some reduction in pain sensitivity has been reported resulting from these transplants, particularly for the reduction of low intensity chronic pain. In most reports, significant differences between control and transplanted animals were noted only after nicotine

administration to stimulate opioid peptide production. However, there have been some reports that analgesia has been observed in a rat chronic pain model from basal level activity of chromaffin tissue allografts.

See, e.g., Vaquero et al., NeuroReport, 2, pp. 149-51 (1991) and Hama and Sagen, Brain Research, 651, pp. 183-93 (1994).

Bovine adrenal chromaffin cells have been encapsulated to form a bioartificial organ ("BAO") for implantation into rats for the treatment of acute and chronic pain. See, e.g., Sagen et al., J. Neurosci., 13, pp. 2415-23 (1993) and Hama et al., 7th World Congress Pain, Abstract 982, Paris France (1993). Initial trials in human subject have been conducted using encapsulated bovine chromaffin cells. See, Aebischer et al., Transplantation, 58, pp. 1275-77 (1994).

There have also been attempts to induce antinociception using other cells, e.g., AtT-20 cells.

20 AtT-20 cells were originally derived from a mouse anterior pituitary tumor. These cells synthesize and secrete \( \beta\)-endorphin. See, e.g., Wu et al., \( \beta\). Neural Transpl. \( \beta\) Plasticity, 5, pp. 15-26 (1993).

AtT-20/hENK cells are AtT-20 cells that have been genetically engineered to carry the entire human proenkephalin A gene (i.e. containing 6 met-enkephalin sequences and one leu-enkephalin sequence) with 200 bases of 5'-flanking sequence and 2.66 kilobases of 3'-flanking sequence. See Wu et al., supra, Comb et al.,

EMBO J., 4, pp. 3115-22 (1985).

Wu et al., J. Neural Transpl. \( \begin{array}{c} \text{Plasticity}, \text{ Plasticity}, \

5, pp. 15-26 (1993) refers to rat hosts transplanted

with AtT-20 or AtT-20/hENK cells. Unstimulated AtT-20/hENK cells produced more antinociception (tail flick test) than produced by AtT-20 implants. In contrast, isoproterenol stimulation produced more antinociception 5 with AtT-20 cells than with AtT-20/hENK cells. Ibid.

In mice hosts, AtT-20 or AtT-20/hENK implants did not affect basal response to thermal nociceptive stimuli. Mice receiving AtT-20 implants developed tolerance to  $\mbox{$\beta$-endorphin}$  and a  $\mbox{$\mu$-opioid}$  agonist 10 (DAMGO). Mice receiving AtT-20/hENK implants developed tolerance to an  $\delta$ -opioid agonist (DPDPE). In response to repeated doses of an  $\mu$  opiate agonist, mice receiving AtT-20/hENK implants developed less tolerance compared to mice receiving AtT-20 cells or controls.

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The antinociceptive effect of isoproterenol treatment appeared equal in mice receiving AtT-20 or AtT-20/hENK cell implants. See, Wu et al., J. Neuroscience, 14, pp. 4806-14 (1994). Wu et al. speculated that one reason for the absence of additional antinociception in mice implanted with enkephalin producing AtT-20/hENK cells may be due to lack of sensitivity of the behavioral assays. possible reason was that met-enkephalin's known antagonist effect on morphine induced antinociception 25 offset the potentiating effect of the single leu-enkephalin, particularly since there are 6 metenkephalin sequences for each leu-enkephalin sequence in pro-enkephalin A.

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#### Summary of the Invention

The present invention provides a cell line that has been genetically engineered to produce at least one analgesic compound from each of the groups consisting of endorphins, enkephalins, and catecholamines. The cell line may be used in the treatment of pain.

There are advantages to using a cell line over the use of primary cells. Expensive and time 10 consuming testing to ensure safety and performance criteria for cells must be performed for individual isolations of primary cells. Less testing is required. of a cell bank. There is no need to isolate primary cells. Output of the desired analgesics may be more stable since the performance of primary cells may be dependent on the age, sex, health or hormonal status of the donor animal. It is also possible to achieve higher output of the desired products, as well as to engineer specifically modified peptides into the cell line. This permits delivery of multiple analgesics simultaneously. Expression of one or more of the analgesics can be regulated (by using a regulatable promoter to drive expression). In addition, for safety, a "suicide" gene can be incorporated into the 25 cell line. Further, for encapsulation purposes proliferating cells have the advantage that they divide to replace dying or dead cells.

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## Brief Description of the Drawing

Figure 1 is a plasmid map of vector pBS-hPOMC-027, pBS-IgSP-hPOMC-028 and pBS-IgSP-hPOMC-ΔACTH-029.

Figure 2 is a plasmid map of vectors pCEP4-hPOMC-030, pCEP4-hPOMC-031, pcDNA3-hPOMC-034 and pcDNA3-hPOMC-035.

Figure 3 is a plasmid map of vectors pCEP4-hPOMC-ΔACTH-032, pCEP4-hPOMC-ΔACTH-033, pcDNA3-hPOMC-ΔACTH-037.

Figure 4 is a plasmid map of vectors pcDNA3-rTH-044, pcDNA3-rTH $\Delta$ -045, and pcDNA3-rTHDKS-075 (also represented as pcDNA3-rTH $\Delta$ KS-075).

Figure 5 is a plasmid map of vectors pcDNA3-15 rTHΔ-IRES-bDBH-088 and pcDNA3-rTHΔKS-IRES-bDBH-076.

Figure 6 is a plasmid map of vector pZeo-Pcmv-rTHAKS-IRES-bDBH-088.

Figure 7 is a plasmid map of vector pBS-Pcmv-rTHAIRES-bDBH-067.

Figure 8 is a plasmid map of vector pBShPOMC-ΔACTH-IRES-rTHΔIRES-bDBH-068.

Figure 9 is a plasmid map of vector pcDNA3hPOMC-ΔACTH-IRES-rTHΔ-IRES-bDBH-069.

Figure 10 is a plasmid map of vector pcDNA3-25 IRES-Zeocin-072.

Figure 11 is a plasmid map of vector pcDNA3hPOMC-ΔACTH-IRES-rTHΔ-IRES-bDBH-IRES-Zeocin-073.

Figure 12 is a plasmid map of vector pcDNA3-hPROA+KS-091.

## Detailed Description of the Invention

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In order that this invention may be more fully understood, the following detailed description is set forth.

Any suitable cell may be transformed with the recombinant DNA molecules of this invention. Among the contemplated cells are chromaffin cells, including conditionally immortalized chromaffin cells such as those described in WO 96/02646, Neuro-2A, PC12, PC12a, 10 SK-N-MC, AtT-20, and RIN cells including RINa and RINb. Preferably the cell has endogenous prohormone convertases and/or dopa decarboxylases.

SK-N-MC cells, a neuroepithelioma cell line, co-expresses several neuropeptides, including 15 enkephalin, cholecystokinin and gastrin-releasing peptide. See, e.g., Verbeeck et al., J. Biol. Chem., 265, pp. 18087-090 (1990). The pro-enkephalin A gene has been expressed in SK-N-MC cells. See, e.g., Folkesson et al., Mol. Brain Res., 3, pp. 147-54 (1988). We prefer AtT-20 and RIN cells, most preferably RIN cells.

RIN cells are a pancreatic endocrine cell line derived from rat. See, e.g., Horellou et al., J. Physiol., 85, pp. 158-70 (1991). RIN cells are 25 known to endogenously produce GABA and ß-endorphin.

Some of the characteristics of various contemplated cells are shown in Table 1.

**Analgesic Substances** 

NE, met-enkephalin

<u>Cells</u>

Chromaffin

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#### Table 1

Other Components

TH, DDC, DBH, PC

	PC12, PC12a	low NE & met-enkephalin	DDC, DBH, PC			
5	AtT-20	β-endorphin	DDC, PC			
	RINa	β-endorphin, GABA	DDC, PC			
	RINb	β-endorphin	DDC, PC			
	Neuro 2A		DDC, DBH, PC			
10	TH = DDC = DβH = PC =	Tyrosine hydroxylase converts tyrosine – I-dopa Dopamine decarboxylase converts I-dopa – dopamine (DA) Dopamine β-Hydroxylase converts DA – norepinephrine (NE) Prohormone Convertases process POMC to β-endorphin and Proenkephalin A (ProA) to met-enkephalin.				
15	AtT20 =	Mouse pituitary corticotroph cell line that endogenously secretes $\beta$ -endorphln via expression of Pro-opiomelanocortin (POMC).				
	RIN =	RIN = Rat insulinoma				

The primary delivery products include at least one each of an endorphin, an enkephalin and a catecholamine.

Neuro 2A = Mouse neuroblastoma

Enkephalins and endorphins are endogenous opioid peptides in humans. These opioid peptides comprise approximately 15 compounds ranging from 5 to 31 amino acids. These compounds bind to and act at least in part via the same μ opioid receptor as morphine, but are chemically unrelated to morphine. In addition, these compounds stimulate other opiate receptors. Yaksh and Malmberg, Textbook of Pain, 3rd Ed. (Eds. P. Wall and R. Melzack), "Central Pharmacology of Nociceptive Transmission," pp. 165-200, 1994 (New York).

The opioid peptides have common chemical properties, but are synthesized in different pathways.

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 $\beta$ -endorphin, the most abundant endorphin, is synthesized as part of a larger precursor molecule, pro-opiomelanocortin ("POMC"). The POMC molecule contains the full sequence of adrenocorticotrophic hormone ("ACTH"),  $\alpha$ -melanocyte-stimulating hormone (" $\alpha$ -MSH"),  $\beta$ -MSH, and  $\beta$ -lipotropin. The POMC precursor molecule also has the potential to generate other endorphins, including  $\alpha$ -endorphin and gamma-endorphin. Processing of the POMC precursor occurs differently within various tissues according to the localization of cleavage enzymes, such as prohormone convertases, within those tissues.

In the pituitary, POMC is cleaved to produce ACTH and \(\beta\)-endorphin, and the ACTH is not further processed. In contrast, in the hypothalamus, ACTH is converted to \(\beta\)-MSH. While different cell types may synthesize the same primary gene product, the final profile of hormone secretion may differ widely.

This invention contemplates use of a DNA

sequence encoding any suitable endorphin that has
analgesic activity. In addition, analogs or fragments
of these endorphins that have analgesic activity are
also contemplated. Thus the endorphin to be produced
by the cells of this invention may be characterized by
amino acid insertions, deletions, substitutions and
modifications at one or more sites in the naturally
occurring amino acid sequence of the desired endorphin.
We prefer conservative modifications and substitutions
(i.e., those having a minimal effect on the secondary
or tertiary structure of the endorphin and on the
analgesic properties of the endorphin). Such
conservative substitutions include those described by

Dayhoff in Atlas of Protein Sequence and Structure, 5, (1978) and by Argos, Embo J., 3, pp. 779-85 (1989).

Techniques for generating such variants of naturally occurring endorphins are well known. For 5 example, codons in the DNA sequence encoding the wild type endorphin may be altered by site specific mutagenesis.

This invention contemplates using a DNA sequence encoding the entire POMC precursor molecule. 10 This embodiment takes advantage of the host cell's cleavage enzymes (i.e., Prohormone convertase 2) to generate a suite of endorphins, some or all of which may have analgesic properties.

This invention also contemplates use of DNA fragments of the POMC gene that encode a particular desired endorphin.

The DNA and amino acid sequence of POMC are well known. Cochet et al., Nature, 297, pp. 335-9 (1982); Takahashi et al., Nucl. Acids Res., 11, pp. 6847-58 (1983).

We prefer a DNA sequence encoding POMC in which the ACTH coding region has been deleted. The preferred endorphin encoded by this construct is ß-endorphin.

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Some enkephalins are synthesized in the adrenal glands as part of a large protein, proenkephalin A, that contains six repeats of the Metenkephalin sequence and one Leu-enkephalin structure. Met-enkephalin, as well as Met-enkephalin-Arg-Phe and 30 Met-enkephalin-Arg-Gly-Leu have significant antinociceptive activity. See, e.g., Sagen et al., Brain Res., 502, pp. 1-10 (1989).

Other enkephalins, i.e., dynorphins and neoendorphins are derived from a distinct molecule, proenkephalin B. Additional "cryptic" peptides are also encoded within the structure of these precursor proteins, and may be released by "pro-hormone-type" cleavage. See, e.g., Harrison's "Principles Of Internal Medicine", 12th Edition, pp. 1168-69 (1991).

This invention contemplates use of a DNA sequence encoding any suitable enkephalin that has analgesic activity. Analogs and active fragments that have analgesic properties are also contemplated. Such analogs or fragments may thus have amino acid insertions, deletions, substitutions at one or more sites in the naturally occurring amino acid sequence.

15 Such variants may be generated as described above.

This invention contemplates use of a DNA sequence encoding a desired enkephalin in its "mature" form. In addition, this invention contemplates using a DNA sequence encoding the entire pro-enkephalin A precursor, or the entire pro-enkephalin B precursor. Further, we also contemplate using DNA encoding a fusion, or fragment of these sequences, that upon expression yields one or more enkephalin-like molecules that have analgesic properties.

25 We prefer use of a DNA sequence encoding the entire pro-enkephalin A precursor molecule. The DNA and amino acid sequence of pro-enkephalin A are well known. Folkesson, supra. This embodiment takes advantage of the host cell's cleavage enzymes, such as prohormone convertase, to generate a suite of enkephalins, some or all of which may have analgesic

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properties. The preferred enkephalin encoded by this construct is Met-enkephalin.

There are three naturally occurring catecholamines which function as neurotransmitters in the central nervous system; norepinephrine ("NE"), epinephrine ("E"), and dopamine. NE is associated with postganglionic sympathetic nerve endings. NE exerts its effects locally in the immediate vicinity of its release.

Catecholamines are synthesized from the amino acid tyrosine, which is sequentially hydroxylated to form dihydroxyphenylalanine (dopa), decarboxylated to form dopamine, and then hydroxylated on the beta position of the side chain by dopamine beta hydroxylase to form NE. Harrison's, <a href="mailto:supra">supra</a>, pp. 380. NE is N-methylated to E by phenylethanolamine-N methyltransferase ("PNMT").

Hydroxylation of tyrosine by tyrosine hydroxylase ("TH") is the rate limiting step in NE synthesis. Regulation of dopa and NE synthesis in the adrenal medulla may be accomplished by changes in the amount and the activity of TH.

In addition, regulation of synthesis of E from NE may occur by changes in the amount and the activity of phenylethanolamine-N-methyltransferase ("PNMT"). PNMT is inducible by glucocorticoids from the adrenal cortex. <u>Ibid</u>.

Catecholamines are maintained in high concentration in adrenal medullary chromaffin tissue, mostly as E. Opioid peptides are also stored in the adrenal gland.

NE and E have similar affinities at  $\alpha_2$ receptors and therefore both potentially contribute to analgesia. Bylund, <u>FASEB J.</u>, 6, PP. 832-39 (1992). The enkephalin peptides that predominantly include met-5 enkephalin selectively activate delta  $(\delta)$  opioid receptors. Reisine and Bell, Trends Neurosci., 16, pp. 506-10 (1993). Activation of  $\alpha_2$  adrenergic and  $\delta$ opioid receptors in the spinal cord each result in antinociception and are potentially synergistic. Yaksh 10 and Malmberg, Progress in Pain Research and Management, Vol. 1, Ed. Fields and Lisbeskind, IASP Press, Seattle, pp. 141-71 (1994). Activation of  $\delta$  versus ( $\mu$ ) opioid receptors in experimental animals results in fewer adverse side effects including constipation and 15 addiction liability (Lee et al., J. Pharmacol. Exp. Ther., 267, pp. 883-87 (1993). The combined delivery of different opioidergic and adrenergic agents may decrease the magnitude of tolerance that develops to a single agent and lead to sustained pain relief. Yaksh and Reddy, Anesthesiol., 54, pp. 451-67 (1981).

This invention contemplates use of a DNA sequence encoding catecholamine biosynthetic enzymes or analogs or fragments thereof to obtain catecholamines that have analgesic properties. The preferred catecholamines in this invention are NE and E.

In one embodiment, the host cell is transformed with the genes necessary to accomplish production of NE or E, as desired. The selection of heterologous gene sequences required depends upon the complement of catecholamine synthesizing enzymes normally occurring in the host cell. For example, RIN cells, and AtT-20 cells lack tyrosine hydroxylase

("TH") and dopamine beta hydroxylase ("DBH"). However, RIN and AtT-20 cells contain endogenous dopa decarboxylase ("DDC"). If the desired catecholamine is E, then the gene encoding PNMT is also required. The gene encoding PNMT is known. Baetge et al., Proc. Nat'l Acad. Sci., 83, pp. 5455-58 (1986).

The gene encoding TH is known. See, e.g., United States patent 5,300,436, incorporated herein by reference. Modified TH variants are also known.

United States patent 5,300,436. In addition, truncated versions of TH that contain the necessary C-terminal catalytic domains are also known. See, e.g., Daubner et al., Protein Science, 2, pp. 1452-60 (1993).

AtT-20 cells have been transformed with wild type TH, as well as various TH muteins. See, e.g., Wu et al., J. Biol. Chem., 267, pp. 25754-758 (1992).

The sequence of the DBH gene is also well known. See, e.g., Lamoroux et al., EMBO J., 6, pp. 3931-37 (1987).

It will be appreciated that in addition to the preferred DNA sequences described herein, there will be many degenerate DNA sequences that code for the desired analgesics.

Secondary compounds with potential analgesic action may also be produced by the cells of this invention. Such compounds include galanin and somatostatin. In addition, neuropeptide Y, neurotensin and cholecystokinin may be produced by the transformed cells of this invention. The cells of this invention may normally produce some or all of these compounds, or may be genetically engineered to do so using standard techniques.

Standard methods may be used to obtain or synthesize the genes encoding the analgesic compounds to be produced by the cells of this invention.

For example, the complete amino acid sequence of the desired compound may be used to construct a back-translated gene. A DNA oligomer containing a nucleotide sequence coding for the desired analgesic compound may be synthesized. For example, several small oligonucleotides coding for portions of each 10 desired polypeptide may be synthesized and then ligated. The individual oligonucleotides typically contain 5' or 3' overhangs for assembly.

The DNA sequence encoding each desired analgesic compound, may or may not also include DNA sequences that encode a signal sequence. Such signal sequence, if present, should be one recognized by the cell chosen for expression of the analgesic compound. It may be prokaryotic, eukaryotic or a combination of the two. It may also be the signal sequence of the 20 native compound. It generally is preferred that a signal sequence be encoded and most preferably that the native signal sequence be used.

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Once assembled, the DNA sequences encoding the desired compounds will be inserted into one or more 25 expression vectors and operatively linked to expression control sequences appropriate for expression in the desired transformed cell.

Proper assembly may be confirmed by nucleotide sequencing, restriction mapping, and 30 expression of a biologically active polypeptide in the transformed cell. As is well known in the art, in order to obtain high expression levels of a transfected

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gene in a host, the gene must be operatively linked to transcriptional and translational expression control sequences that are functional in the chosen expression cell.

The choice of expression control sequence and expression vector will depend upon the choice of cell. A wide variety of expression host/vector combinations may be employed. Useful expression vectors for eukaryotic hosts, include, for example, vectors 10 comprising expression control sequences from SV40, bovine papilloma virus, adenovirus and cytomegalovirus.

We prefer pcDNA3, pCEP4, pZeoSV (InVitrogen, San Diego) and pNUT.

Any of a wide variety of expression control sequences may be used in these vectors. Such useful expression control sequences include the expression control sequences associated with structural genes of the foregoing expression vectors. Examples of useful expression control sequences include, for example, the 20 early and late promoters of SV40 or adenovirus, the promoter for 3-phosphoglycerate kinase or other glycolytic enzymes, the promoters of acid phosphatase, e.g., Pho5, the promoters of the yeast  $\alpha$ -mating system and other sequences known to control the expression of 25 genes of eukaryotic cells or their viruses, and various combinations thereof.

It should of course be understood that not all vectors and expression control sequences will function equally well to express the DNA sequences described herein. Neither will all cells function equally well with the same expression system. However, one of skill in the art may make a selection among

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these vectors, expression control sequences and cells without undue experimentation. For example, in selecting a vector, the host cell must be considered because the vector must replicate in it. The vector's copy number, the ability to control that copy number, and the expression of any other proteins encoded by the vector, such as antibiotic markers, should also be considered.

In selecting an expression control sequence,
a variety of factors should also be considered. These
include, for example, the relative strength of the
sequence, its controllability, and its compatibility
with the actual DNA sequence encoding the desired
analgesic compounds, particularly as regards potential
secondary structures. Host cells should be selected by
consideration of their compatibility with the chosen
vector, the toxicity of the product coded for by the
DNA sequences, their secretion characteristics, their
ability to fold the polypeptides correctly, and their
culture requirements. If the host cell is to be
encapsulated, cell viability when encapsulated and
implanted in a recipient should also be considered.

Within these parameters, one of skill in the art may select various vector/expression control sequence/host combinations that will express the desired DNA sequences in culture.

In one embodiment, cells (e.g., RIN cells) are sequentially transformed with 4 separate expression vectors containing the POMC gene, the pro-enkephalin A gene, the TH gene and the DBH gene. In such a transformed host cell, amplification of copy number of the heterologous genes is more difficult to achieve.

Thus use of fewer expression vectors is preferred. Most preferably, a single expression vector, containing all 4 heterologous genes, is used.

In a particular embodiment RIN cells are 5 sequentially transformed with 3 expression vectors. The first vector contains the POMC gene operably linked to the CMV promoter. Preferably a truncated version of the POMC gene is used, having the ACTH coding region deleted. The second vector contains the pro-enkephalin 10 A gene operably linked to the CMV promoter. Preferably the proA construct contains the Kozak sequence ' immediately upstream of the start codon. The third vector contains both the TH gene (preferably truncated and having the Kozak consensus sequence immediately 15 upstream of the start codon) and the DBH gene. In this embodiment, the TH gene is operably linked to the CMV promoter. The DBH gene is operably linked to an internal ribosome entry site promoter sequence. RIN cells are then transformed sequentially with each expression vector according to known protocols.

In another embodiment, a single expression vector containing the pro-enkephalin A gene, the POMC gene, the TH gene, and the DBH gene is constructed.

Preferably, the ACTH region of the POMC gene is deleted. Preferably the TH gene is truncated.

Multiple gene expression from a single transcript is preferred over expression from multiple transcription units. One approach for achieving expression of multiple genes from a single eukaryotic transcript takes advantage of sequences in picorna viral mRNAs known as internal ribosome entry sites ("IRES"). These sites function to facilitate protein

translation from sequences located downstream from the first AUG of the mRNA.

Macejak and Sarnow reported that the 5' untranslated sequence of the immunoglobulin heavy chain binding protein (BiP, also known as CRP 78, the glucose-regulated protein of molecular weight 78,000) mRNA can directly confer internal ribosome binding to an mRNA in mammalian cells, in a 5'-cap independent manner, indicating that translation initiation by an internal ribosome binding mechanism is used by this cellular mRNA. Nature 353, pp. 90-94 (1991).

WO 94/24870 refers to use of more than two IRES for translation initiation from a single transcript, as well as to use of multiple copies of the same IRES in a single construct.

"suicide" gene in the transformed cells. Most preferably, the cell carries the TK (thymidine kinase) gene as a safety measure, permitting the host cell to be killed in vivo by treatment with gancyclovir.

Use of a "suicide" gene is known in the art.

See, e.g., Anderson, published PCT application

WO 93/10218; Hamre, published PCT application

WO 93/02556. The recipient's own immune system

25 provides a first level of protection from adverse reactions to the implanted cells. If encapsulated, the polymer capsule itself may be immuno-isolatory. The presence of the TK gene (or other suicide gene) in the expression construct adds an additional level of safety

30 to the recipient of the implanted cells.

Preferred vectors for use in this invention include those that allow the DNA encoding the analgesic

compounds to be amplified in copy number. Such amplifiable vectors are well known in the art. They include, for example, vectors able to be amplified by DHFR amplification (see, e.g., Kaufman, United States Patent 4,470,461, Kaufman and Sharp, "Construction Of A Modular Dihydrafolate Reductase cDNA Gene: Analysis Of Signals Utilized For Efficient Expression", Mol. Cell. Biol., 2, pp. 1304-19 (1982)) or glutamine synthetase ("GS") amplification (see, e.g., United States patent 5,122,464 and European published application 338,841). Such amplification can be used to increase output of the desired analgesic compounds.

Other techniques for increasing the output of the desired analgesic compounds are contemplated. For example, subcloning existing polyclonal cell lines is contemplated. Cells are cloned by limiting dilution to a single cell in each well. Cell clones are cultures, and the clones are tested to select the clone with the highest output of analgesic substances.

Another technique for increasing the output of the desired analgesic compounds involves cloning altered forms of biosynthetic enzymes with higher activity than the wild type form (i.e., the truncated TH 1-155). Some truncated forms of TH have 4-6 times increased activity over the wild type form of TH. See, e.g., Daubner et al., "Expression and characterization of catalytic and regulatory domains of rat tyrosine hydroxylase" Protein Science, 2, pp. 1452-60 (1993).

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In addition, use of tyrosine-free media to select to increase tetrahydrobiopterin cofactor levels may potentially increase tyrosine hydroxylase activity. See, e.g., Horellou et al., "Retroviral transfer of a

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human tyrosine hydroxylase cDNA in various cell lines; regulated release of dopamine in mouse anterior pituitary AtT-20 cells", Proc. Natl. Acad. Sci. USA, 86, pp. 7233-37 (1989).

Preferably, the output of B-endorphin ranges between 1 and 10,000 pg/106 cells/hr. Preferably, the output of met-enkephalin ranges between 1 and 10,000 pg/106 cells/hr. Preferably, the output of catecholamines ranges between 1 and 1,000 pmoles/106 10 cells/hr.

The cells of this invention may be implanted into a mammal, including a human, for the treatment of pain. If implanted unencapsulated, any suitable implantation protocol may be used, including those 15 outlined by Sagen et al., United States patent 4,753,635, incorporated herein by reference.

It may be desirable to encapsulate the genetically modified cells of this invention before implantation. Such encapsulated cells form a 20 bioartificial organ ("BAO"). BAOs may be designed for implantation in a recipient or can be made to function extra-corporeally. The BAOs useful in this invention typically have at least one semipermeable outer surface membrane or jacket surrounding a cell-containing core.

25 The jacket permits the diffusion of nutrients, biologically active molecules and other selected products through the BAO. The BAO is biocompatible.

In some cases, the membrane may serve to also immunoisolate the cells by blocking the cellular and molecular effectors of immunological rejection. use of immunoisolatory membranes allows for the implantation of allo and xenogeneic cells into an

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individual without the use of immunosuppression. If biologically active molecules are released from the isolated cells, they pass through the surrounding semipermeable membrane into the recipient's body. If metabolic functions are provided by the isolated cells, the substances to be metabolized enter the BAO from the recipient's body through the membrane to be acted on by the cells.

A variety of types of membranes have been 10 used in the construction of BAOs. Generally, the membranes used in BAOs are either microporous of ultrafiltration grade membranes. A variety of membrane materials have been suggested for use in BAOs, including PAN/PVC, polyurethanes, polysufones, 15 polyvinylidienes, and polystyrenes. Typical membrane geometries include flat sheets, which may be fabricated into "sandwich" type constructions, having a layer of living cells positioned between two essentially planar membranes with seals formed around the perimeter of the 20 device. Alternatively, hollow fiber devices may be used, where the living cells are located in the interior of a tubular membrane. Hollow fiber BAOs may be formed step-wise by loading living cells in the lumen of the hollow fiber and providing seals on the 25 ends of the fiber. Hollow fiber BAOs may also be formed by a coextrusion process, where living cells are coextruded with a polymeric solution which forms a membrane around the cells.

BAOs have been described, for example, in United States patent Nos. 4,892,538, 5,106,627, 5,156,844, 5,158,881, and 5,182,111, and PCT Application Nos. PCT/US/94/07015, WO 92/19195, WO

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93/03901, and WO 91/00119, all of which are incorporated herein by reference.

BAOs may contain other components that promote long term survival of the encapsulated cells.

5 For example, WO 92/19195 refers to implantable immunoisolatory biocompatible vehicles having a hydrogel matrix for enhancing cell viability.

The encapsulating membrane of the BAO may be made of a material which is the same as that of the core, or it may be made of a different material. In either case, a surrounding or peripheral membrane region of the BAO which is permselective and biocompatible will be formed. The membrane may also be constructed to be immunoisolatory, if desired. The core contains isolated cells, either suspended in a liquid medium or immobilized within a hydrogel matrix.

The choice of materials used to construct the BAO is determined by a number of factors and is described in detail in Dionne WO 92/19195. Briefly, various polymers and polymer blends can be used to manufacture the capsule jacket. Polymeric membranes forming the BAO and the growth surfaces therein may include polyacrylates (including acrylic copolymers), polyvinylidenes, polyvinyl chloride copolymers, polyurethanes, polystyrenes, polyamides, cellulose acetates, cellulose nitrates, polysulfones, polyphosphazenes, polyacrylonitriles, poly(acrylonitrile/covinyl chloride), as well as derivatives, copolymers and mixtures thereof.

BAOs may be formed by any suitable method known in the art. One such method involves coextrusion of a polymeric casting solution and a coagulant which

can include biological tissue fragments, organelles, or suspensions of cells and/or other therapeutic agents, as described in Dionne, WO 92/19195 and United States Patents 5,158,881, 5,283,187 and 5,284,761,

5 incorporated herein by reference.

The jacket may have a single skin or a double skin. A single-skinned hollow fiber may be produced by quenching only one of the surfaces of the polymer solution as it is co-extruded. A double-skinned hollow fiber may be produced by quenching both surfaces of the polymer solution as it is co-extruded.

Numerous capsule configurations, such as cylindrical, disk-shaped or spherical are possible.

The jacket of the BAO will have a pore size 15 that determines the nominal molecular weight cut off (nMWCO) of the permselective membrane. Molecules larger than the nMWCO are physically impeded from traversing the membrane. Nominal molecular weight cut off is defined as 90% rejection under convective 20 conditions. In situations where it is desirable that the BAO is immunoisolatory, the membrane pore size is chosen to permit the particular factors being produced by the cells to diffuse out of the vehicle, but to exclude the entry of host immune response factors into 25 the BAO. Typically the nMWCO ranges between 50 and 200 kD, preferably between 90 and 150 kD. The most suitable membrane composition will also minimize reactivity between host immune effector molecules known to be present at the selected implantation site, and 30 the BAO's outer membrane components.

The core of the BAO is constructed to provide a suitable local environment for the particular cells

isolated therein. The core can comprise a liquid medium sufficient to maintain cell growth. Liquid cores are particularly suitable for maintaining transformed cell lines like PC12 cells. Alternatively, the core can comprise a gel matrix. The gel matrix may be composed of hydrogel (alginate, "Vitrogen", etc.) or extracellular matrix components. See, e.g., Dionne WO 92/19195.

three general classes. The first class carries a net negative charge (e.g., alginate). The second class carries a net positive charge (e.g., collagen and laminin). Examples of commercially available extracellular matrix components include Matrigel™ and Vitrogen™. The third class is net neutral in charge (e.g., highly crosslinked polyethylene oxide, or polyvinylalcohol).

Any suitable method of sealing the BAO may be used, including the employment of polymer adhesives and/or crimping, knotting and heat sealing. These sealing techniques are known in the art. In addition, any suitable "dry" sealing method can also be used. In such methods, a substantially non-porous fitting is provided through which the cell-containing solution is introduced. Subsequent to filling, the BAO is sealed. Such a method is described in copending United States application Serial No. 08/082,407, herein incorporated by reference.

One or more in vitro assays are preferably
used to establish functionality of the BAO prior to
implantation in vivo. Assays or diagnostic tests well
known in the art can be used for these purposes. See,

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e.g., Methods In Enzymology, Abelson [Ed], Academic Press, 1993. For example, an ELISA (enzyme-linked immunosorbent assay), chromatographic or enzymatic assay, or bioassay specific for the secreted product can be used. If desired, secretory function of an implant can be monitored over time by collecting appropriate samples (e.g., serum) from the recipient and assaying them. If the recipient is a primate, microdialysis may be used.

The number of BAOs and BAO size should be sufficient to produce a therapeutic effect upon implantation is determined by the amount of biological activity required for the particular application. In the case of secretory cells releasing therapeutic substances, standard dosage considerations and criteria known to the art are used to determine the amount of secretory substance required. Factors to be considered are discussed in Dionne, WO 92/19195.

Implantation of the BAO is performed under
sterile conditions. Generally, the BAO is implanted at
a site in the host which will allow appropriate
delivery of the secreted product or function to the
host and of nutrients to the encapsulated cells or
tissue, and will also allow access to the BAO for
retrieval and/or replacement. The preferred host is a
primate, most preferably a human.

A number of different implantation sites are contemplated. These implantation sites include the central nervous system, including the brain, spinal cord, and aqueous and vitreous humors of the eye. Preferred sites in the brain include the striatum, the cerebral cortex, subthalamic nuclei and nucleus Basalis

of Meynert. Other preferred sites are the cerebrospinal fluid, most preferably the subarachnoid space and the lateral ventricles. This invention also contemplates implantation into the kidney subcapsular site, and intraperitoneal and subcutaneous sites, or any other therapeutically beneficial site.

In order that this invention may be better understood, the following examples are set forth.

These examples are for purposes of illustration only, and are not to be construed as limiting the scope of this invention in any manner.

#### Examples

## Construction of Polycistronic Expression Vectors

## Construction of IgSP-POMC Fusion

The Smal-Sall fragment containing the human POMC exon 3 was subcloned into pBS cloning vector (Stratagene). See <u>Takahashi</u>, <u>supra</u>; <u>Cochet</u>, <u>supra</u>. The resulting plasmid was named as pBS-hPOMC-027. See Fig. 1.

20 A PCR fragment was generated using two oligonucleotide primers, termed oCNTF-003 (SEQ ID NO: 1) and oIgSP-018, (SEQ ID NO: 2) and the pNUT plasmid containing the human CNTF gene. See Baetge et al., <a href="Proc. Natl. Acad. Sci. USA">Proc. Natl. Acad. Sci. USA</a>, 83, pp. 5454-58 (1986). Both primers oCNTF-003 and oIgSP-018, contain synthetic BamHI and SmaI restriction sites, respectively, at the 5' ends.

The 196 base pair (bp) PCR fragment was digested with restriction endonucleases BamHI and the SmaI-isoschizomer XmaI, and electrophoresed through an

1% SeaPlaque agarose. The 193 bp HindIII/XmaI DNA fragment was excised and purified using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME).

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pBS-hPOMc-027 was also digested with BamHI and XmaI and purified from 1% SeaPlaque agarose using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME). The ligation mixture was transformed into E. coli DH5 $\alpha$  (Gibco BRL, Gaithersburg, MD).

Positive sub-clones were initially identified by the cracking gel procedure (Promega Protocols and Applications Guide, 1991). Minilysate DNA was then prepared using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME) and subject to BamHI and SmaI restriction digestions. The positive sub-clone was named as pBS-IgSP-hPOMC-028. See Fig. 1. The nucleotide sequence of the fusion junction in pBS-IgSP-hPOMC-028 was determined by the dideoxynucleotide sequence determination using the Sequenase kit (USBC, Cleveland). The sequence of the IgSP-hPOMC fusion is shown in SEQ ID NO: 3.

# Construction of IgSP-POMC Expression Vectors

The IgSP-hPOMC DNA fragment in pBS-IgSP-hPOMC-028 was subcloned into pcDNA3 (Invitrogen Corp., San Diego, CA) and pCEP4 (Invitrogen Corp., San Diego, CA) in sense and anti-sense orientations.

The NotI-SalI IgSP-hPOMC fragment from pBS-IgSP-hPOMC-028 was ligated with the NotI-XhoI digested pCEP4 resulting in the sense orientation clone named as pCEP4-hPOMC-030. Fig. 2. The BamHI-SalI IgSP-hPOMC fragment from pBS-IgSP-hPOMC-028 was ligated with the

BamHI-XhoI digested pCEP4 resulting in the anti-sense orientation clone named as pCEP4-hPOMC-031. Fig. 2. The insert orientation in pCEP4-hPOMC-030 and -031 was confirmed by BamHI, NotI, SalI and NotI/SalI restriction digestions as well as by dideoxynucleotide sequence determination using the Sequenase kit (USBC, Cleveland).

The BamHI-SalI IgSP-hPOMC fragment from pBS-IgSP-hPOMC-028 was ligated with the BamHI-XhoI digested pcDNA3 resulting in the sense orientation clone named as pcDNA3-hPOMC-034. Fig. 2. The NotI-HindIII IgSP-hPOMC fragment from pBS-IgSP-hPOMC-028 was ligated with the NotI-HindIII digested pcDNA3 resulting in the antisense orientation clone named as pcDNA3-hPOMC-035.

Fig. 2. Restriction digestion using SmaI, BamHI, EcoRI, and BamHI/EcoRI was used to confirm the insert orientation in pcDNA3-hPOMC-034, whereas HindIII, NotI and SalI were used for pcDNA3-hPOMC-035.

## Construction of ACTH Deleted IgSP-POMC

The ACTH coding region in the POMC gene in pBS-IgSP-hPOMC-028 was deleted. pBS-IgSP-hPOMC-028 was first digested with XmaI restriction enzyme and treated with pfu DNA polymerase (Promega, Madison, WI). The XmaI-pfu DNA polymerase treated pBS-IgSP-hPOMC-028 was then digested with StuI restriction enzyme and purified from 1% SeaPlaque agarose using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME). The self-ligation mixture was transformed into E. coli DH5α (Gibco BRL, Gaithersburg, MD). Positive sub-clones were identified by BamHI/HindIII restriction digestion and named as pBS-IgSP-hPOMCΔACTH-029. See Fig. 1. The

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nucleotide sequence of the ACTH deletion region in pBS-IgSP-hPOMC-ΔACTH-029 was confirmed by the dideoxynucleotide sequence determination. The sequence of the IgSP-hPOMC-ΔACTH fusion is shown in SEQ ID 5 NO: 4.

# Construction of ACTH Deleted IgSP-POMC Expression Vectors

The IgSP-hPOMC-ΔACTH DNA fragment in pBS-IgSP-hPOMC-ΔACTH-029 was subcloned into pcDNA3 10 (Invitrogen Corp., San Diego, CA) and pCEP4 (Invitrogen Corp., San Diego, CA) in sense and anti-sense orientations. The NotI-SalI IgSP-hPOMC-ΔACTH fragment from pBS-IgSP-hPOMC-ΔACTH-029 was ligated with the NotI-XhoI digested pCEP4 resulting in the sense orientation clone named as pCEP4-hPOMC-ΔACTH-032 (Fig. 3). The BamHI-SalI IgSP-hPOMC-ΔACTH fragment from pBS-IgSP-hPOMC-ΔACTH-029 was ligated with the BamHI-XhoI digested pCEP4 resulting in the anti-sense orientation clone named as pCEP4-hPOMC-ΔACTH-033 (Fig. 3). The insert orientation in pCEP4-hPOMC-ΔACTH-032 and -033 was confirmed by BamHI and EcoRI restriction digestions as well as by dideoxynucleotide sequence determination using the Sequenase kit (USBC, Cleveland).

The BamHI-SalI IgSP-hPOMC-ΔACTH fragment from pBS-IgSP-hPOMC-ΔACTH-029 was ligated with the BamHI-XhoI digested pcDNA3 resulting in the sense orientation clone named as pcDNA3-hPOMΔACTH-036 (Fig. 3). The NotI-HindIII IgSP-hPOMC-ΔACTH fragment from pBS-IgSP-30 hPOMC-ΔACTH-029 was ligated with the NotI-HindIII

digested pcDNA3 resulting in the anti-sense orientation clone named as pcDNA3-hPOMC- $\Delta$ ACTH-037 (Fig. 3).

Restriction digestion using PvuII and EcoRI was used to confirm the insert orientation in pcDNA3-5 hPOMC-ΔACTH-036, whereas SalI and EcoRI were used for pcDNA3-hPOMC-ΔACTH-037.

# Cloning of Full Length and Truncated TH cDNA

Total RNA from PC12 cells was prepared using the guanidinium thiocyanate-based TRI reagent (Molecular Research Center, Inc., Cincinnati, OH). 10 Five hundred ng of PC12 total RNA was reverse transcribed at 42°C for 30 minutes in a 20µl reaction volume containing 10 mM Tris.HC1 (pH 8.3), 50 mM KC1, 4 mM of each dNTP, 5 mM MgCl $_2$ , 1.25  $\mu$ M oligo (dT) 15mer, 1.25 µM random hexamers, 31 units of RNase Guard RNase Inhibitor (Pharmacia, Sweden) and 200 units of SuperScript II reverse transcriptase (Gibco BRL, Gaithersburg, MD). Two micro-liters of the above reverse transcribed cDNA was added to a 25 µl PCR 20 reaction mixture containing 10 mM Tris.HCl (pH 8.3), 50 mM KC1, 800 of each nM dNTP, 2 mM MgC12, 400 nM of primers #1 and #2, and 2.5 units of Thermus aquaticus (Taq) DNA polymerase (Boehringer Mannheim, Germany).

To generate the full length TH cDNA,

25 oligonucleotide primers orTH-052 (SEQ ID NO: 5) and

orTH-053 (SEQ ID NO: 6) were used. For the truncated

TH, primers orTH-054 (SEQ ID NO: 7) and orTH-053 (SEQ

ID NO: 6) were used instead. These oligonucleotides

were constructed based on published TH sequence

30 information in Grima et al., Nature, 326, pp. 707-11

(1987); US patent 5,300,436, and Daubner, supra.

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Gaithersburg, MD).

Primers orTH-052 (SEQ ID NO: 5) and orTH-054 (SEQ ID NO: 7) have synthetic HindIII restriction site at the 5' end where orTH-053 has BamHI at the 5' end. The PCR reaction mixtures were subject to 30 amplification cycles consisted of: denaturation, 94°C 30 seconds (first cycle 2 minutes); annealing, 50°C 1 minute; and extension, 72°C 3.5 minutes (last cycle 5 minutes). The 1537 bp full length and 1087 bp truncated rat TH PCR fragments were digested with restriction endonucleases BamHI and HindIII and resolved on an 1% SeaPlaque agarose gel. The 1531-bp and 1081-bp HindIII/BamHI DNA fragments were excised and purified using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME).

pcDNA3 expression vector was also digested with BamHI and HindIII and purified from 1% SeaPlaque agarose using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME). The ligation mixture was transformed into E.coli DH5 $\alpha$  (Gibco BRL,

Cracking gel procedure (Promega Protocols and Applications Guide, 1991) was used to screen out the positive sub-clones. The identity of the correct clones was further verified by BamHI/HindIII double digestion.

The positive sub-clones for the full-length and truncated rat TH in pcDNA3 were named as pcDNA3-rTH-044 (Fig. 4) and pcDNA3-rTHA-045 (Fig. 4), respectively. The nucleotide sequence of both full-length and truncated rat TH PCR clones was determined by the dideoxynucleotide sequence determination using

the Sequenase kit (USBC, Cleveland). The sequence of the rTH $\Delta$  construct is shown in SEQ ID NO: 16.

To optimize the translation efficiency of the truncated rat TH, oligonucleotide primer orTH-078 (SEQ ID NO: 8) was designed so that the consensus Kozak sequence is immediate up stream to the start codon ATG. pcDNA3-rTHA-45 was used as the template in a 50 µl PCR reaction mixture with reagent composition identical to the one described above with the exception that the oligonucleotide primers were replaced with orTH-078 (SEQ ID NO: 8) and orTH-053 (SEQ ID NO: 6). The 1097 bp PCR product was cloned into pcDNA3 in the same manner as described above. The resulting sub-clone was named pcDNA3-rTHAKS-75 (Fig 4). The sequence of the rTHAKS construct is shown in SEQ ID NO: 17.

## Construction of rTH-IRES-bDBH Fusion Gene

Recombinant PCR methodology was used to generate the rTH-IRES-bDBH fusion gene.
Oligonucleotides oIRES-057 (SEQ ID NO: 9) and obDBH-065
20 (SEQ ID NO: 10) are specific for IRES and bDBH gene sequences, respectively, and contain synthetic BamHI and NotI restriction sites at the 5' end, respectively.
Oligonucleotides oIRES-bDBH-064 (SEQ ID NO: 11) and oIRES-bDBH-066 (SEQ ID NO: 12) are complementary to each other. Furthermore, oligonucleotide primer oIRES-bDBH-064 (SEQ ID NO: 11) has its 5' 16 nucleotides identical to the IRES sequence and its 3' 18 nucleotides identical to the bDBH sequence; and vice versa for oIRES-bDBH-066 (SEQ ID NO: 12).

Two first PCR reactions were carried out using oligonucleotide pairs oIRES-057/oIRES-bDBH-066

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and oIRES-bDBH-064/obDBH-065 on templates pCTI-001
(with an insert containing the IRES sequence shown in SEQ ID NO: 30) and pBS-bDBH-006 (containing the bovine DBH gene cloned from bovine adrenal chromaffin cells,

5 Lamoroux et al., EMBO J., 6, pp. 3931-37 (1987))
plasmids, respectively. One hundred ng of template DNA was added to a 50 µl PCR reaction mixture containing
10 mM Tris.HCl (pH 8.3), 50 mM KCl, 800 of each nM dNTP, 2 mM MgCl2, 400 nM of primers #1 and #2, and 2.5

10 units of Thermus aquaticus (Taq) DNA polymerase
(Boehringer Mannheim, German).

The PCR reaction mixtures were subject to 30 amplification cycles consisted of: denaturation, 94 °C for 30 seconds (first cycle 2 minutes); annealing, 15 50 °C 1 minute; and extension, 72 °C 30 seconds (last cycle 5 minutes). The PCR products were resolved on 1% TrivieGel 500 (TrivieGen). Two agarose plugs containing each one of the first PCR products were transfer to a tube containing 50 µl of PCR reaction 20 mixtures identical to the one described above with the exception that the oligonucleotides oIRES-057 and obDBH-065 were used.

The second PCR reaction was subject to 30 amplification cycles consisted of: denaturation, 94 °C for 30 seconds (first cycle 2 minutes); annealing, 60 °C 30 seconds (second to fourth cycles 37 °C 2 minutes); and extension, 72 °C 30 seconds (last cycle 2 minutes). The 2407 bp IRES-bDBH fusion PCR product and the cloning vector pcDNA3-rTHA-45 were digested with BamHI and NotI restriction enzymes and subsequently purified from 1% SeaPlaque agarose gel using the FMC

SpinBind DNA purification kit (FMC BioProducts, Rockland, ME).

The ligation of IRES-bDBH/BamHI/Notl and pcDNA3-rTHA-045/BamHI/NotI would generate a rTHA-IRES-5 bDBH expression vector named as pcDNA3-rTHΔ-IRES-bDBH-066 (Fig. 5) whereas that of IRES-bDBH/BamHI/NotI and pcDNA3-rTHAKS-075/BamHI/NotI would generate a rTHAKS-IRES-bDBH expression vector, named as pcDNA3-rTHAKS-IRES-bDBH-076 (Fig. 5), where the start codon ATG in 10 rTHA is preceded with a consensus Kozak sequence. The sequence of the rTHA-IRES-bDBH construct is shown in SEQ ID NO: 18. The sequence of the rTHAKS-IRES-bDBH construct is shown in SEQ ID NO: 19. The ligation mixture was transformed into DH5α (Gibco BRL, 15 Gaithersburg, MD). The positive clones were identified by the cracking gel procedure (Promega, Madison, WI) and restriction digestions using HindIII, BamHI, HindIII/BamHI, SmaI and NotI.

The 4114 bp NruI-XhoI fragment containing the CMV promoter-rTHAKS-IRES-bDBH was excised out of pcDNA3-rTHAKS-IRES-bDBH-076 and subcloned into pZeoSV cloning vector (Invitrogen Corp., San Diego, CA) digested with ScaI and XhoI in the multiple cloning site. The resulting expression vector was named as pZeo-Pcmv-rTHAKS-IRES-bDBH-088 (Fig. 6).

# Construction of IgSP-hPOMC ACTHrTHD-IRES-bDBH Fusion Gene

The 4100 bp NruI-NotI fragment containing the CMV promoter, rTHD-IRES-bDBH fusion gene, and BGH polyadenylation sequence was excised out of pcDNA3-

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rTHA-IRES-bDBH-066 and subcloned into the pBS (Stratagene, La Jolla, CA) cloning vector.

The resulting plasmid pBS-Pcmv-rTHA-IRES-bDBH-067 (Fig. 7) was used as the intermediary construct to which the recombinant PCR IgSP-hPOMCDACTH-IRES fragment would be inserted.

Oligonucleotide oIgSP-068 (SEQ ID NO: 13), containing a synthetic EcoRV restriction site, is specific for the IgSP sequence.

Oligonucleotide primer orTHA-073 (SEQ ID NO: 14) is specific for the rTHA sequence and contains an endogenous Smal restriction site.

Oligonucleotide primers ohPOMC-IRES-069 (SEQ ID NO: 15) and ohPOMC-IRES-070 (SEQ ID NO: 20) are complementary to each other. Furthermore, oligonucleotide primer ohPOMC-IRES-069 has its 5', 18 nucleotides identical to the hPOMC sequence and its 3' 12 nucleotides identical to the IRES sequence; and vice versa for ohPOMC-IRES-070.

Oligonucleotide primers oIRES-rTHΔ-071 (SEQ ID NO: 21) and oRIRES-rTHΔ-072 (SEQ ID NO: 22) are complementary to each other. In addition, oligonucleotide primer oIRES-rTHΔ-071 has its 5' 15 nucleotides identical to the rTHΔ sequence and its 3' 18 nucleotide identical to the IRES sequence; and vice versa for oRIRES-rTHΔ-072.

Three sets of first PCR reactions were carried out.

PCR reaction A: template pBS-IgSP-hPOMCDACTH-029, oligonucleotides oTgSP-068/ohPOMC-IRES-069;

PCR reaction B: template pCTI-001, oligonucleotides ohPOMC-IRES-070/oIRES-rTHΔ-071; and

PCR reaction C: template pcDNA3-rTH $\Delta$ -045, oligonucleotides orIRES-rTH $\Delta$ -072/orTH $\Delta$ -073.

The three sets of first PCR reactions were carried in 50 µl PCR reaction mixture containing 100 ng of template DNA, 10 mM Tris. HCl (pH 8.3), 50 mM KCl, 800 of each nM dNTP, 2 mM MgCl23, 400nM of primers #l and #2, and 2.5 units of Thermus aquaticus (Taq) DNA polymerase (Boehringer Mannheim, Germany).

The PCR reaction mixtures were subject to 30 amplification cycles consisted of: denaturation, 94 °C for 30 seconds (first cycle 2 minutes); annealing, 50 °C 1 minute; and extension, 72 °C 30 seconds (last cycle 5 minutes).

The PCR products were resolved on 1%

TrivieGel 500 (TrivieGen). Two agarose plugs containing each one of the PCR products from PCR reactions B and C were transferred to a tube containing 50 µl of PCR reaction mixtures identical to the one described above with the exception that the oligonucleotides ohPOMC-IRES-070 and orTHA-073 were used.

The second PCR reaction was subject to 30 amplification cycles consisted of: denaturation, 94 °C for 30 seconds (first cycle 2 minutes); annealing, 60 °C 30 seconds (second to fourth cycles 37 °C 2 minutes); and extension, 72 °C 30 seconds (last cycle 2 minutes).

The PCR products were treated as described above. Agarose plugs containing the PCR products from the second PCR reaction and the PCR reaction A were combined and subjected to a third PCR amplification using oIgSP-068/rTHA-073. The 1203 bp IgSP-hPOMC-IRES-

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rTHΔ fusion PCR product and the cloning vector pBS-PCmv-rTHΔ-IRES-bDBH-067 were digested with EcoRV and XmaI restriction enzymes and subsequently purified from 1% SeaPlaque agarose gel using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME). The ligation mixture was transformed into DH5α (Gibco BRL, Gaithersburg, MD).

The positive clones were identified by the cracking gel procedure (Promega, Madison, WI) and restriction digestions using EcoRI, KpnI and NotI. The resulting clone was named as pBS-IgSP-hPOMCAACTH-IRES-rTHA-IRES-bDBH-068. Fig. 8. The sequence of this construct is shown in SEQ ID NO: 23.

# Construction of IgSP-hPOMCACTH-IRESrTHA-IRES-bDBH Expression Vectors

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The 4491 bp NotI fragment containing the IgSP-hPOMCΔACTH-IRES-rTHΔ-IRES-bDBH gene was excised out of the pBS-IgSP-hPOMCΔACTH-IRES-rTHΔ-IRES-bDBH-068 and subcloned into the pcDNA3 (Invitrogen Corp., San Diego, CA) at the NotI site in the multiple cloning site. Restriction digestion using NotI and SmaI confirmed that the IgSP-hPOMCΔACTH-IRES-rTHΔ-IRES-bDBH gene was inserted in the sense orientation resulting in pcDNA3-IgSP-hPOMCΔACTH-IRES-rTHΔ-IRES-bDBH-069. See Fig. 9.

# Construction of IgSP-hPOMCAACTH-IRES-rTHA-IRES-bDBH-IRES-Zeocine Expression Vector

Recombinant PCR methodology was used to generate the IRES-Zeocine fusion gene.

30 Oligonucleotides oIRES-074 (SEQ ID NO: 24) and oZeocin-

077 (SEQ ID NO: 25) are specific for IRES and Zeocin gene sequences, respectively, and contain synthetic NotI and XhoI restriction sites at the 5' end, respectively. Oligonucleotides oIRES-Zeocin-075 (SEQ ID NO: 26) and oIRES-Zeocin-076 (SEQ ID NO: 27) are complementary to each other. Furthermore, oligonucleotide oIRES-Zeocin-075 has its 5'15 nucleotides identical to the Zeocin sequence and its 3' 18 nucleotides identical to the IRES sequence; and vice versa for oIRES-Zeocin-076.

Two first PCR reactions were carried out using oligonucleotide pairs oIRES-074/oIRES-Zeocin-075 and oIRES-Zeocin-076/oZeocin-075 on templates pCTI-001 and pZeoSV (Invitrogen Corp., San Diego, CA) plasmids, respectively.

One hundred ng of template DNA was added to a 50 µl PCR reaction mixture containing 10mM Tris.HCl (pH 8.3), 50 mM KCl, 800 of each nM dNTP, 2 mM MgCl2, 400 nM of primers #1 and #2, and 2.5 units of Thermus aguaticus (Taq) DNA polymerase (Boehringer Mannheim, Germany).

The PCR reaction mixtures were subject to 30 amplification cycles consisted of: denaturation, 94 °C for 30 seconds (first cycle 2 minutes); annealing, 25 50 °C 1 minute; and extension, 72 °C 30 seconds (last cycle 5 minutes).

The PCR products were resolved on 1%
TrivieGel 500 (TrivieGen). Two agarose plugs
containing each one of the first PCR products were
transfer to a tube containing 50 µl of PCR reaction
mixtures identical to the one described above with the

exception that the oligonucleotides oIRES-074 and oZeocin-077 were used.

The second PCR reaction was subject to 30 amplification cycles consisted of: denaturation, 94 °C for 30 seconds (first cycle 2 minutes); annealing, 50 °C 30 seconds (second to fourth cycles 37 °C 2 minutes); and extension, 72 °C 30 seconds (last cycle 2 minutes).

The 974 bp IRES-Zeocin fusion PCR product and the cloning vector pcDNA3 were digested with NotI and XhoI restriction enzymes and subsequently purified from 1% SeaPlaque agarose gel using the FMC SpinBind DNA purification kit (FMC BioProducts, Rockland, ME).

The ligation of IRES-Zeocin/NotI/XhoI and pcDNA3/NotI/XhoI would generate an intermediate cloning vector named as pcDNA3-IRES-Zeocin-072. Fig. 10.

The positive clones were identified by the cracking gel procedure (Promega, Madison, WI) and restriction digestions using HindIII, SmaI, XhoI, NotI and NotI/XhoI.

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To generate the final IgSP-hPOMCDACTH-IRESrTHD-IRES-bDBH-IRES-Zeocine Expression Vector, a 4491 bp NotI fragment containing the IgSP-hPOMCΔACTH-IRESrTHΔ-IRES-bDBH gene was excised out of the pBS-IgSP-25 hPOMCΔACTH-IRES-rTHΔ-IRES-bDBH-068 (Fig. 8; SEQ ID NO: 23) and subcloned in to the pcDNA3-IRES-Zeocin-072 (Fig. 10) at the NotI site in the multiple cloning site.

Restriction digestion using NotI and SmaI confirmed that the IgSP-hPOMCAACTH-IRES-rTHA-IRES-bDBH gene was inserted in the sense orientation resulting in pcDNA3-IgSP-hPOMCAACTH-IRES-rTHA-IRES-bDBH-IRES-Zeocin-

073. The sequence of this construct is shown in SEQ ID NO: 28. Fig. 11.

### Construction of ProA+KS Fusion

A construct containing the coding region of
the human pro-enkephalin A gene with the consensus
Kozak sequence immediately upstream to the start codon
ATG. The sequence of this construct is shown in SEQ ID
NO: 29.

# Construction of hProA+KS Expression Vector

The HindIII/BamHI fragment containing the hProA+KS fusion was ligated into BamHI and Hind III digested pcDNA3 expression vector substantially as described above. After screening as described above, a positive sub-clone was named pcDNA3-hProA+KS-091.

Fig. 12. Construction of the pBS-CMV Pro A vector is detailed in Mothis, J. and Lindberg, I., Endocrinology, 131, pp. 2287-96 (1992).

#### Transformation of Cells

RIN and AtT-20 cells were transformed as follows.

The RINa and AtT-20 based cell lines were grown in DMEM (Gibco) with 10% fetal bovine serum and pen-strep-fungizone (Gibco) base media. The cells were plated out in P100 petri dishes (750,000 cells/dish) in 10 ml of base media. 18-24 hours later, the cells were transfected using calcium phosphate method with a kit made by Stratagene (San Diego, CA). A 10 µg amount of the plasmid vector DNA was diluted in 450 µl of deionized sterile water. Then, 50 µl of a 10x buffer

(solution #1) was added to the plasmid DNA. A 500 µl amount of solution #2 was immediately added to the DNA containing solution and mixed gently. This was incubated at room temperature for 20 minutes and then the 1.0 ml solution was added to the cells in the petri dish. The cells were incubated overnight and 18-24 hours later the cells were washed 2x with Hanks balanced salt solution without calcium and magnesium. Then, the cells were cultured in base media + selection drugs. The cells were selected in either 600 µg/ml geneticin (Gibco) or 400 µg/ml hygromycin (Boehringer Mannheim) or 500 µg/ml Zeocin (In Vitrogen, San Diego, CA). Cells were sequentially transfected and selected to obtain the final cell line.

The RINa cells were transfected with plasmid pCEP4-hPOMC-030 containing the POMC gene. This is a hygromycin resistant vector. The cells were also transformed with plasmid pcDNA3-hProA+KS-091. This is a geneticin resistant vector. Finally, the cells were transfected with plasmid pZeo-PCMV-rTHΔKS-IRES-bDBH-088 which conferred Zeocin resistance.

The AtT-20 cells were transfected with plasmid pBS-CMV-ProA and pCEP4-POMC-ΔACTH-32 which conferred geneticin and hygromycin resistance, respectively. Finally, the cells were transfected with plasmid pZeo-Pcmv-rTHΔKS-IRES-bDBH-088.

We have tested a number of media for cell growth. Surprisingly we have found that in certain serum-free medias, the above cell lines have enhanced neurotransmitter output, compared to serum-containing media. We prefer CHO-Ultra (Biowhitaker) for the

growth of AtT-20 cells, and Ultra-Culture (Biowhitaker) for the growth of RINa cells.

Output of various analgesics from one transformed RINa cell line (RINa/ProA/P030/P088) is 5 shown in Table 2. All values represent unstimulated cells. Output of ß-endorphin and met-enkephalin is in pg/10<sup>6</sup> cells/hr. ß-endorphin and met-enkephalin were measured by radioimmunoassay using Incstar kits (Stillwater, Minnesota). Catecholamine output is in 10 pmoles/10 cells/hr. The numbers in parentheses represent values from cells that were preincubated 18 hours with 100 µM tetrahydrobiopterin. Catecholamines were measured by high performance liquid chromatography as described in Lavoie et al., "Two PC12 15 pheochromocytoma lines sealed in hollow fiber-based capsules tonically release 1-dopa in vitro", Cell transplantation, 2, pp. 163-73 (1993). GABA output from these RINa cells was 28 ng/10<sup>6</sup> cells/hrs.

## Table 2

20	<u>Cell Line</u>	Endogenous Analgesic Substances	<u>B-endorphin</u>	<u>Met-enk</u>	<u>DA</u> E
25	RIN a/ ProA/ POMC/ TH-IRES-DBH	β-endorphin GABA	22	17	3 0 (6) (2)

There are encrypted enkephalin fragments which are not fully processed from the pro-enkephalin precursor molecule. These encrypted enkephalins have opioid receptor binding activity. We digested these encrypted enkephalins to measure opioid activity. The trypsin digest protocol is as follows. A 2 µg/ml trypsin (Worthington #34E470) solution is added to media

samples on ice. Samples are vortexed, then incubated for 20 minutes in a 37°C waterbath. After the 20 minute digest, samples are returned to ice and 100 ng/ml carboxypeptidase B (Sigma #C-7011) is added. 5 Samples are mixed by vortexing, and returned to the 37°C waterbath for 15 minutes. Samples are placed on ice once more and 10 ug/ml trypsin inhibitor is added. At this stage, samples are either extracted for metenkephalin or immediately frozen for future extraction. 10 This results in the full enzymatic cleavage to free all met-enkaphalin from the longer encrypted fragments. A met-enkaphalin radioimmunoassay of the digested sample gives total met-enkaphalin from the supermatant. The transformed RINa cells appear to have greater than 5 fold more encrypted enkaphalins compared to fully processed met-enkaphalin.

# Fiber capsule formation and characteristics

Hollow fibers are spun from a 12.5-13.5% poly(acrylonitrile vinylchloride) solution by a wet spinning technique. Cabasso, Hollow Fiber Membranes, vol. 12, Kirk-Othmer Encyclopedia of Chemical Technology, Wiley, New York, 3rd Ed. pp. 492-517 (1980), Unites States patent 5,158,881, incorporated herein by reference.

The resulting membrane fibers may either be double skinned or single skinned PAN/PVC fibers. In order to make implantable capsules, lengths of fiber are first cut into 5 cm long segments and the distal extremity of each segment sealed with an acrylic glue. 30 Encapsulation hub assemblies are prepared by providing lengths of the membrane described above, sealing one

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end of the fiber with a single drop of LCM 24 (Light curable acrylate glue, available from ICI), curing the glue with blue light, and repeating the step with a second drop. The opposite end is previously attached 5 to a frangible necked hub assembly, having a silicone septum through which the cell solution may be introduced. The fiber is glued to the hub assembly by applying LCM 22 to the outer diameter of the hub assembly, pulling the fiber up over it, and curing with 10 blue light. The hub/fiber assemblies are placed in sterilization bags and are ETO sterilized.

Following sterilization with ethylene oxide and outgassing, the fibers are deglycerinated by ultrafiltering first 70% EtOH, and then HEPES buffered 15 saline solution through the walls of the fiber under vacuum.

# Preparation and Encapsulation of Transformed Cells

The transformed cells are prepared and encapsulated as follows:

20

A matrix solution is prepared using a commercially available alginate, collagen or other suitable matrix material. The cell solution was diluted in the ratio of two parts matrix solution to one part cell solution containing the transformed cells 25 described above. We prefer Vitrogen (Celtix, Santa Clara) as a matrix for AtT-20 cells.

We prefer Organogen (Organogenesis, Canton, MA) as a matrix for RINa cells. The RINa based cells are prepared for encapsulation by the following method. 30 The cells are grown in base media of DMEM + 10% fetal bovine serum during the proliferation phase. These

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cells can be removed from the tissue culture flasks by two washes in Hanks balanced salt solution without calcium and magnesium. Then the cells are incubated in 0.25% trypsin + EDTA for 1 minute. This is removed and the cells are rinsed free of the flask using Hanks balanced salt solution without calcium and magnesium solution. The cells are placed in 10 mls of base media and centrifuged at 100 x g for 2 minutes. The cells are resuspended in 10 mls of the preferred serum free media (Ultra culture, Biowhitaker, Walkersville, MD). Surprisingly, the RINa cells secrete more analgesic substances when cultured in this serum free media relative to serum continuing base media.

The cells are centrifuged at 100 g twice in
the preferred serum free media before the cells are
concentrated 1:1 with the preferred Organogen matrix.
Organogen is a 1% bovine tendon collagen obtained as a
sterile solution. 8 parts of this solution are mixed
with 1 part 10X DPBS. 0.5 N sodium hydroxide is added
until physiological pH is attained (approximately
250 µls).

The final concentration of the cell + matrix solution used for encapsulation can range from 20,000 - 50,000 cells/µl. The cells are counted in a standard manner on a hemocytometer.

The cell/matrix suspension is placed in a 1 ml syringe. A Hamilton 1800 Series 50 microliter syringe is set for a 15 microliter air bubble, is inserted into a 1 ml syringe containing the cell solution and 30 microliters are drawn up. The cell solution is injected through the silicone seal of the hub/fiber assembly into the lumen of a modacrylic

hollow fiber membrane with a molecular weight cutoff of approximately 50,000-100,000 daltons. Ultrafiltration should be observed along the entire length of the fiber. After one minute, the hub is snapped off the sub-hub, exposing a fresh surface, unwet by cell solution. A single drop of LCM 24 is applied and the adhesive cured with blue light. The device is placed first in HEPES buffered NaCl solution and then in CaCl<sub>2</sub> solution for five minutes to cross-link the alginate.

10 Each implant is about 5 cm long, 1 mm in diameter, and contained approximately 2.5 million cells.

After the devices are filled and sealed, a silicone tether (Speciality Silcone Fabrication, Paso Robles, CA) (ID: 0.69, OD: 1.25) is then placed over the proximal end of the fiber. A radiopaque titanium plug is inserted in the lumen of the silicone tether to act as a radiographic marker. The devices are then placed in 100 mm tissue culture dishes in 1.5 ml PC-1 medium, and stored at 37°C, in a 5% CO<sub>2</sub> incubator for in vitro analysis and for storage until implantation.

The encapsulated cells are then implanted into the human sub-arachnoid space as follows:

## Surgical Procedure

After establishing IV access and

administering prophylactic antibiotics (cefazolin sodium, 1 gram IV), the patient is positioned on the operating table, generally in either the lateral decubitus or genu-pectoral position, with the lumbar spine flexed anteriorly. The operative field is

sterily prepared and draped exposing the midline dorsal lumbar region from the levels of S-1 to L-1, and

allowing for intraoperative imaging of the lumbar spine with C-arm fluoroscopy. Local infiltration with 1.0% lidocaine is used to establish anesthesia of the skin as well as the periosteum and other deep connective tissue structures down to and including the ligamentum flavum.

A 3-5 cm skin incision is made in the parasagital plane 1-2 cm to the right or left of the midline and is continued down to the lumbodorsal 10 fascia using electrocautery for hemostasis. Using traditional bony landmarks including the iliac crests and the lumbar spinous processes, as well as fluoroscopic guidance, and 18 gauge Touhy needle is introduced into the subarachnoid space between L-3 and 15 L-4 via an oblique paramedian approach. The needle is directed so that it enters the space at a shallow, superiorly directed angle that is no greater than 30-35° with respect to the spinal cord in either the sagittal or transverse plane. Appropriate position of 20 the tip of the needle is confirmed by withdrawal of several ml of cerebrospinal fluid (CSF) for preimplantation catecholamine, enkephalin, glucose, and protein levels and cell counts.

that the opening at the tip is oriented superiorly
(opening direction is marked by the indexing notch for
the obturator on the needle hub), and the guide wire is
passed down the lumen of the needle until it extends 45 cm into the subarachnoid space (determined by
premeasuring). Care is taken during passage of the
wire that there is not resistance to advancement of the
wire out of the needle and that the patient does not

complain of significant neurogenic symptoms, either of which observations might indicate misdirection of the guide wire and possible impending nerve root or spinal cord injury.

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After the guide wire appears to be appropriately placed in the subarachnoid space, the Touhy needle is separately withdrawn and removed from the wire. The position of the wire in the midline of the spinal canal, anterior to the expected location of the caud equina, and without kinks or unexplainable 10 bends is then confirmed with fluoroscopy. After removal of the Touhy needle the guide wire should be able to be moved freely into and out of the space with only very slight resistance due to the rough surface of 15 the wire running through the dense and fibrous ligamentum flavum.

The 7 French dilator is then placed over the guide wire and the wire is used to direct the dilator as it is gently but firmly pushed through the fascia, paraspinous muscle, and ligamentum flavum, following the track of the wire toward the subarachnoid space. Advancement of the 7 French dilator is stopped and the dilator removed from the wire as soon as a loss of resistance is detected after passing the ligamentum flavum. This is done in order to avoid advancing and manipulating this relatively rigid dilator within the subarachnoid space to any significant degree.

After the wire track is "overdilated" by the 7 French dilator, the 6 French dilator and cannula 30 sheath are assembled and placed over the guide wire. The 6 French dilator and cannula are advanced carefully into the subarachnoid space until the opening tip of

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the cannula is positioned 7 cm within the space. As with the 7 French dilator, the assembled 6 French dilator and cannula are directed by the wire within the lumen of the dilator. Position within the subarachnoid space is determined by premeasuring the device and is grossly confirmed by fluoroscopy. Great care is taken with manipulation of the dilators and cannula within the subarachnoid space to avoid misdirection and possible neurologic injury.

When appropriate positioning of the cannula is assured, the guide wire and the 6 French dilator are gently removed from the lumen of the cannula in sequence. Depending on the patient's position on the operating table, CSF flow through the cannula at this 15 point should be noticeable and may be very brisk, requiring capping the cannula or very prompt placement of the capsule implant in order to prevent excessive CSF.

10

The encapsulated (transformed cells) is 20 provided in a sterile, double envelope container, bathed in transport medium, and fully assembled including a tubular silicone tether. Prior to implantation through the cannula and into the subarachnoid space, the capsule is transferred to the insertion kit tray where it is positioned in a location that allowed the capsule to be maintained in transport medium while it is grossly examined for damage or major defects, and while the silicone tether is trimmed, adjusting its length to the pusher and removing the hemaclip™ that plugs its external end.

The tether portion of the capsule is mounted onto the stainless steel pusher by inserting the small WO 96/40959 PCT/US96/09629

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diameter wire portion of the pusher as the membrane portion of the device is carefully introduced into the cannula. The capsule is advanced until the tip of the membrane reaches a point that is 2-10 mm within the cranial tip of the cannula in the subarachnoid space. This placement is achieved by premeasuring the cannula and the capsule-tether-pusher assembly, and it assures that the membrane portion of the capsule is protected by the cannula for the entire time that it is being advanced into position.

After the capsule is positioned within the cannula, the pusher is used to hold the capsule in position (without advancing or withdrawing) in the subarachnoid space while the cannula is completely 15 withdrawn from over the capsule and pusher. The pusher is then removed from the capsule by sliding its wire portion out of the silicone tether. Using this method the final placement of the capsule is such that the 5 cm long membrane portion of the device lay entirely 20 within the CSF containing subarachnoid space ventral to the cauda equina. It is anchored at its caudal end by a roughly 1-2 cm length of silicone tether that runs within the subarachnoid space before the tether exits through the dura and ligamentum flavum. The tether continues externally from this level through the paraspinous muscle and emerges from the lumbodorsal fascia leaving generally 10-12 cm of free tether material that is available for securing the device.

CSF leakage is minimized by injecting fibrin
glue (Tissel®) into the track occupied by the tether in
the paraspinous muscle, and by firmly closing the
superficial fascial opening of the track with a purse-

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string suture. The free end of the tether is then anchored with non-absorbable suture and completely covered with a 2 layer closure of the skin and subcutaneous tissue.

The patient is then transferred to the neurosurgical recovery area and kept at strict bed rest, recumbent, for 24 hours postoperatively.

Antibiotic prophylaxis is also continued for 24 hours following the implantation procedure.

## 10 Sequences

The following is a summary of the sequences set forth in the Sequence Listing:

- SEQ ID NO:1 -- DNA sequence of oligo oCNTF-003
- SEQ ID NO:2 -- DNA sequence of oligo oIgSP-018
- 15 SEQ ID NO:3 -- DNA sequence of IgSP-hPOMC fusion
  - SEQ ID NO:4 -- DNA sequence of IgSP-hPOMC-ΔACTH fusion
  - SEQ ID NO:5 -- DNA sequence of oligo orTH-052
  - SEQ ID NO:6 -- DNA sequence of oligo orTH-053
  - SEQ ID NO:7 -- DNA sequence of oligo orTH-054
- 20 SEQ ID NO:8 -- DNA sequence of oligo orTH-078
  - SEQ ID NO:9 -- DNA sequence of oligo oIRES-057
  - SEQ ID NO:10 -- DNA sequence of oligo obDBH-065
  - SEQ ID NO:12 -- DNA sequence of oligo oIRES-bDBH-066

SEQ ID NO:11 -- DNA sequence of oligo oIRES-bDBH-064

- 25 SEQ ID NO:13 -- DNA sequence of oligo oIRE-068
  - SEQ ID NO:14 -- DNA sequence of oligo orTH $\Delta$ -073
  - SEQ ID NO:15 -- DNA sequence of oligo ohPOMC-IRES-069
  - SEQ ID NO:16 -- DNA sequence of rTHA1-155
  - SEQ ID NO:17 -- DNA sequence of rTHA+KS
- 30 SEQ ID NO:18 -- DNA sequence of rTHΔ-IRES-bDBH
  - SEQ ID NO:19 -- DNA sequence of rTHAKS-IRES-bDBH

SEQ ID NO:20 -- DNA sequence of oligo ohPOMC-IRES-070
SEQ ID NO:21 -- DNA sequence of oligo oIRES-rTHA-071
SEQ ID NO:22 -- DNA sequence of oligo orIRES-rTHA-072
SEQ ID NO:23 -- DNA sequence of IgSP-hPOMCAACTH-IRESrTHA-IRES-bDBH-068 fusion
SEQ ID NO:24 -- DNA sequence oIRES-074
SEQ ID NO:25 -- DNA sequence of oligo oZeocin-077
SEQ ID NO:26 -- DNA sequence of oligo oIRES-Zeocin-075
SEQ ID NO:27 -- DNA sequence of oligo oIRES-Zeocin-076
SEQ ID NO:28 -- DNA sequence IgSP-hPOMCAACTH-IRES-rTHA
-IRES-bDBH-IRES-Zeocin-073
SEQ ID NO:29 -- DNA sequence of proA+KS

SEQ ID NO:30 -- DNA sequence of IRES fragment

# Deposits

5

15 RINa/ProA/POMC/TH-IRES-DBH cells, transformed to produce a catecholamine, an enkephalin and an endorphin, as described above in the example (and in Table 2), named RINa/ProA/PO30/PO88, have been deposited. The deposit was made in accordance with the Budapest Treaty and was deposited at the American Type Culture Collection, Rockville, Maryland, U.S.A. on June 7, 1995. The deposit received accession number CRL 11921.

The foregoing description has been for the

purpose of illustration and description only. This

description is not intended to limit the invention to

the precise form exemplified. It is intended that the

scope of the invention be defined by the claims

appended hereto.

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# SEQUENCE LISTING

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	(2) INFO	RMATION FOR SEQ ID NO:1:	
10	(i)	SEQUENCE CHARACTERISTICS:  (A) IENGIH: 33 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
••	(ii)	MOLECULE TYPE: CINA	
	(iii)	HYPOIHETICAL; NO	
15	(iv)	ANTI-SENSE: NO	···
20	(vii)	IMEDIATE SOURCE: (B) CLONE: contr-003	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:1:	
25	COCCATION	OG OGTCACCOCT AGAGTOGAGC TGT	33
25	(2) INFO	RMATION FOR SEQ ID NO:2:	
30	·(i)	SEQUENCE CHARACTERISTICS:  (A) LENGIH: 23 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
35	(ii)	MOLECULE TYPE: CINA	
	(iii)	HYPOIHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
40	(vii)	IMMEDIATE SOURCE: (B) CLONE: 01gSP-018	
45	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:2:	
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5 (A) (E) (C) (I) (I) (I) (I) (I)	JENCE CHARACTERISTICS:  A) IENGIH: 849 base pairs  B) TYPE: nucleic acid  C) STRANIFINESS: single  D) TOPOLOGY: linear  FOULE TYPE: INA (genomic)  FOULETICAL: NO  TI-SENSE: NO  FEDIATE SOURCE:
	COIHEITICAL: NO TI-SENSE: NO TEDIATE SOURCE:
(iii) HYF	TI-SENSE: NO FDIATE SOURCE:
\	EDIATE SOURCE:
(iv) AN	
(vii) IM	3) CLONE: IgSP-hPCMC
•	ATURE: A) NAME/KEY: 5'UTR B) LOCATION: 143
	ATURE:  A) NAME/KEY: excon  B) LOCATION: 4489
•	ATURE: A) NAME/KEY: intron B) LOCATION: 90168
· ·	ATURE: A) NAME/KEY: 3'UTR B) LOCATION: 807849
C	ATURE:  A) NAME/KEY: misc_feature  B) LOCATION: 43186  D) OTHER INFORMATION: /product= "IgSp region"
40	
Ċ	ATURE:  A) NAME/KEY: misc_feature  B) LOCATION: 187806  D) OTHER INFORMATION: /product= "hPOMC region
45	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

(2) INFORMATION FOR SEQ ID NO:3:

	CENTOCCCITY CACCOCITY ACTUAL CITCHACTICIT CACCOCITY ACAMICAAAT COACCIGGET	60
5	TATCTICITC CIGATGECAG TEGITACAGG TAAGGGGCCC CCAAGTCCCA AACITGAGGG	120
	TOTATIANACT CIGIGACAGT GECANICACT TIGOCITICT TICIACAGG GIGANTICGG	180
	CITIOCOGG ANATIGGORAC GAGCAGOCIC TGACOGAGAA COCCOGGAAG TACGTCATGG	240
10	GOCACTICOG CIGGGACCGA TICCGCCCCC CCAACAGCAG CAGCAGCGGC AGCAGCGGC	300
	CAGGGCAGAA GOGGGAGGAC GICICAGGGG GOGAAGACIG GGGGAGGGGG GCIGAGGGG	360
	COUNTAGE CONTAGE CONTA	. 420
15	CCATGEAGCA CITICOGCTGG GGCAAGCOGG TGGGCAAGAA GCGGGGCCCA GTGAAGGTGT	480
	ACCUTANCES CECCEPAGEAC GAGICOGOCC AGRICUTOCC CCTGEAGUTC AAGAGGGAGC	540
20	TEACTECOCA COCACTOCOC CACCEACATG COCOCCACT CACCECCAG	600
	GEGOTTAGET CEPACTIGEAR CACAGOTTICE TEGTIGGOGGE CEPACAPAGA GACAPAGGOC	660
25	CCTACAGGAT GGAGCACTIC CGCTGGGGCA GCCCCCAA GGACAAGCGC TACGGCGGTT	720
	TCATGACCIC CCAGAAGAGC CAGACGCCCC TGGTGACGCT GTTCAAAAAAC GCCATCATCA	780
	AGANCECCIA CANGANGEEC GAGTGAGGEC ACAGGGGCCCC CCAGGGCTAC CCTCCCCCAG	840
30	GAGGIOGAC	849

# (2) INFORMATION FOR SEQ ID NO:4:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGIH: 525 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic) 40

(iii) HYPOIHETICAL: NO

(iv) ANTI-SENSE: NO

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(vii) IMEDIATE SOURCE:

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	(B) CLONE: Igsp-hpomodacih	
5	(ix) FEATURE:  (A) NAME/KEY: 5'UIR  (B) LOCATION: 143	
	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 4489	
10	(ix) FEATURE: (A) NAME/KEY: intron (B) LOCATION: 90168	
15	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 169482	•
20	(ix) FEATURE: (A) NAME/KEY: 3'UIR (B) LOCATION: 483525	
25	(ix) FFATURE:  (A) NAME/KEY: misc feature  (B) LOCATION: 44188  (D) OTHER INFORMATION: /product= "Igsp region"	
30	(ix) FEATURE:  (A) NAME/KEY: misc feature  (B) LOCATION: 189482  (D) OTHER INFORMATION: /product= "hPOMC region"	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:  GEATOGGGT CACCCIAGA GIOGAGCIGT GAOGGICCIT ACAAIGAAAT GCAGCIGGGT	60
	TATCTICTIC CIGATGGCAG TGGITACAGG TAAGGGGCIC CCAAGICCCA AACTIGAGG	120
40	TOCATAAACT CIGIGACAGT GECAATCACT TIGOCITICT TICIACAGEG GIGAATTOEG	180
	CTTICCCCCC CAGTICAAGA GGGAGCTGAC TGGCCAGCGA CTCCGGGAGG	240
45	CACATICECCO COCATICACI COCCATICACI COCCAGGOCCAC CTICCACCACA	300
7.5	GOCIGCIGGI GGOGGOGGAG AAGAAGGACG AGGGOCCCIA CAGGAIGGAG CACTIOOGCI	36

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	GCCCAGCCC GCCCAAAGGAC AAGCCCIACG GCCGITTICAT GACCICCCAG AAGAGCCAGA	420
	CECCCCIGGI CACCCIGITIC AAAAACCCCA TCATCAAGAA CCCCTACAAG AAGGCCAGT	480
5	GAGGGCACAG CGGGCCCACCCTC CCCCAGGAGG TOGAC	525
	(2) INFORMATION FOR SEQ ID NO:5:	
10	(i) SEQUENCE CHARACTERISTICS:  (A) LENGIH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
15	(ii) MOIFOULE TYPE: cDNA	
	(iii) HYPOIHETICAL: NO	•
20	(iv) ANTI-SENSE: NO	
	(vii) IMMEDIATE SOURCE: (B) CLONE: orTH-052	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
	COCAAGCTTG CACTATGOOC ACCOCAGUG	30
30	(2) INFORMATION FOR SEQ ID NO:6:	
35	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CINA	
40	(iii) HYPOIHEITCAL: NO	
	(iv) ANTI-SENSE: NO	
45	(vii) IMMEDIATE SOURCE: (B) CLONE: orTH-053	

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	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
	CCCGGATCCT ATGCATTIAG CTAATGGCAC	30
5	(2) INFORMATION FOR SEQ ID NO:7:	
10	(i) SEQUENCE CHARACTERISTICS:  (A) LENGIH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDELNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CINA	
15	(iii) HYPOIHETICAL: NO	
	(iv) ANTI-SENSE: NO	: ` · · · · · · · · · · · · · · · · · ·
20	(vii) IMMEDIATE SOURCE: (B) CLONE: orTH-054	•
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
	COCAAGCITA TEGICOCCIG GITCOCAAGA	30
	(2) INFORMATION FOR SEQ ID NO:8:	
30	(i) SEQUENCE CHARACTERISTICS:  (A) IENGIH: 33 base pairs  (B) TYPE: nucleic acid	
35	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
	(iii) HYPOTHETICAL: NO	
40	(iv) ANTI-SENSE: NO	
	(vii) IMMEDIATE SOURCE:	

(B) CLONE: orTH-078

45

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
	COCAMICTIC GOCACCATEG TOCCTEGIT CCC	33
5	(2) INFORMATION FOR SEQ ID NO:9:	
10	(i) SEQUENCE CHARACIERISTICS:  (A) LENGIH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDELNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CINA	
15	(iii) HYPOIHETICAL: NO	
	(iv) ANTI-SENSE: NO	
20	(vii) IMEDIATE SOURCE: (B) CLONE: oIRES-057	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
23	ANAGENTOG COCCICTOCC TOCCOCCC	30
	(2) INFORMATION FOR SEQ ID NO:10:	
30	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid	
	(C) STRANDEINESS: single (D) TOPOLOGY: linear	
35	(ii) MOLECULE TYPE: CDNA	
	(iii) HYPOIHETICAL: NO	
40	(iv) ANTI-SENSE: NO	
45	(vii) IMMEDIATE SOURCE: (B) CLONE: obDBH-065	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

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	AAAGCCCCCC CCCACGITICA GCCTTTGCCC	30
	(2) INFORMATION FOR SEQ ID NO:11:	
10	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	
	(iii) HYPOTHETICAL: NO	
15	(iv) ANTI-SENSE: NO	
20	(vii) IMMEDIATE SCURCE: (B) CLONE: oIRES-bDBH-064	•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
25	CITIGOCACAA OCATIGIACEG CACUGOGGIG	30
	(2) INFORMATION FOR SEQ ID NO:12:	
30	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: single  (D) TOPOLOGY: linear	•
35	(ii) MOLECULE TYPE: CLNA	
	(iii) HYPOIHETICAL: NO	
40	(iv) ANTI-SENSE: NO	
	(VII) IMEDIATE SOURCE: (B) CLONE: OIRES-blbH-066	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	

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	OGCIGGIGCOG TACATGGITG TGGCAAGCIT	30
	(2) INFORMATION FOR SEQ ID NO:13:	
5	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: CLNA	
	(iii) HYPOTHETICAL: NO	
15	(iv) ANTI-SENSE: NO	
20	(vii) IMMEDITATE SOURCE: (B) CLONE: 0IgSP-068	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
25	AAAGATATOG OGGOOGIC ACCOCTAGAG	30
23	(2) INFORMATION FOR SEQ ID NO:14:	
30	(i) SEQUENCE CHARACTERISTICS:  (A) IENGIH: 25 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: cDNA	
35	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
40	(vii) IMMEDIATE SOURCE: (B) CLONE: orTHD-073	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
	ATACACCIEG TCAGAGAAGC COGGG	25

ATACACCIGG TCAGAGAGC CCCGG

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	(2) INFO	MATION FOR SEQ ID NO:15:
5	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGIH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
10	(ii)	MOLECULE TYPE: CDNA
	(iii)	HYPOTHETICAL: NO
15	(iv)	ANTI-SENSE: NO
٠	(vii)	IMEDIATE SOURCE: (B) CLONE: ohPOMC-IRES-069
20	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:15:
		GG ACAGGGGCCC GCTGTGCCCT
25		RMATION FOR SEQ ID NO:16:
30		SEQUENCE CHARACTERISTICS:  (A) IENGIH: 1030 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: DNA (genomic)
35	(iii)	HYPOTHETICAL: NO
	(iv)	ANTI-SENSE: NO
40	(vii)	IMMEDIATE SCURCE: (B) CLONE: rTHD
45	(ix)	FEATURE: (A) NAME/KEY: 5'UIR (B) LOCATION: 16
	121	ELENT IDE.

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(A) NAME/KEY: exon
(B) LOCATION: 7..1017

(ix) FEATURE:

5

(A) NAME/KEY: 3'UIR

(B) LOCATION: 1018..1030

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

10 AAGCTTATEG TOOCCIGGIT COCAAGAAAA GIGICGGAAT TOGACAAGIG TCACCACCIG 60 GICACCAAGI TIGACCCIGA ICIGGACCIG GACCACCCGG GCITCICIGA CCAGGIGIAT 120 CECCAGOGIC GEAGCIGAT TECAGAGATT GCCTTCCAGT ACAAGCACGG TGAACCAATT 180 240 CCCCATGIGG AATACACACC GGAAGAGAIT GCIACCIGGA AGGAGGIATA TGICACGCIG AAGGCCTCT ATGCTACCCA TGCCTGCCGG GAGCACCTGG AGGGTTTCCA GCTTCTGGAA - 300 20 COGTACTIGIG CCTACCEAGA CEACAGCATC CCACAGCTGG AGGACGTGTC CCCCTTCTTG 360 420 AAGGAGOGGA CIGGCTIOCA GCIGOGACOC GIGGOOGGIC TACIGIOCGC COGIGATTIT 25 CIGGORAGIC TOCOCTIONS OSTIGITICAA TOCACCAGT ATATOOGOCA TOCCICA 480 CCIATECATT CACCIGAGOC GGACIGCIGC CATGAGCIGT TGGGACATGT ACCCATGTIG 540 GCIGACCECA CATTIECCCA GITICICCCAG GACATIGGAC TIGCATCICT GGGGGCCTCA 600 30 CATCAACAAA TICAAAAACT CICCACGGIG TACIGGITCA CIGIGGAAIT CGGGCIAIGI 660 ANACAGAATG GGGAGCTGAA GGCTTATGGT GCAGGGCTGC TGTCTTCCTA CGGAGAGCTC 35 CIGCACIOCC TGICAGAGA GOCIGAGGIC OGAGOCITIG ACCAGACAC AGCAGCIGIG 780 840 CAGCCCTACC AAGATCAAAC CIACCAGCCT GIGIACTTIG TGICCGAGAG CITCAATGAC GCCAAGGACA AGCICAGGAA CIAIGCCICT CGIATCCAGC GCCCATTCTC TGIGAAGITT 900 40 GACCOGIACA CACIGOCAT TGACGIACIG GACAGOCCIC ACACCATOCA GOGCICCITG 960 CAGGGGGTOC ACCATGAGT COACACCTG COCCAGGCAC TGAGTGCCAT TAGCTAAATG 1020 1030 45 CATAGGATOC

(2) INFORMATION FOR SEQ ID NO:17:

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5	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 1037 base pairs  (B) TYPE: nucleic acid  (C) STRANDELNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLFOULE TYPE: DNA (genomic)	
10	(iii) HYPOIHETICAL: NO	
	(iv) Anti-Sense: No	
15	(vii) IMMEDIATE SOURCE: (B) CLONE: rTHIKS	•
20	(ix) FEATURE: (A) NAME/KEY: 5'UTR (B) LOCATION: 113	
	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 141024	
25	(ix) FEATURE:  (A) NAME/KEY: 3'UIR  (B) LOCATION: 10251037	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
	AAGCITUGOC ACCATGGICC OCTGGITTOOC AAGAAAAGTG TOGGAATTIGG ACAAGTGICA	60
35	CACCIGGIC ACCAAGITIG ACCCIGATOT GGACCIGGAC CACCCGGGCT TOTOTIGACCA	120
	GGIGIATOGC CAGOGIOGGA AGCIGATIGC AGAGATIGOC TICCAGIACA AGCACGIGA	180
40	ACCAPATICOC CATGIGGAAT ACACAGOGGA AGAGATIGCT ACCIGGAAGG AGGIATATGT	240
40	CACECIGAAG GEOCICIATIG CIACOCATIC CIGOOGGEAG CACCIGGAGG GITTOCAGCT	300
	TCIGGAACGG TACIGIGGCT ACCGAGAGGA CAGCATCCCA CAGCIGGAGG ACGIGICCCG	360
45	CITCHIGAAG GAGUGACUG GCITUCAGCT GUGACUGIIG GOOGIICIAC TGICUGUUG	420
	TGATTTTCTG GOCAGTCTGG CCTTCCGCGT GTTTCAATGC ACCCAGTATA TCCGCCATGC	480

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	CIOCICACCI AIGCATICAC CIGAGOOGGA CIGCIGOCAT GAGCIGITEG GACAIGIACC	540
	CATGITIGGCT CACCICACAT TIGGCCAGTT CICCCAGGAC ATTIGGACTIG CATCICTGGG	600
5	GCCICAGAT GAAGAAATIG AAAAACICIC CACGGIGIAC TGGITCACIG TGGAATICGG	660
	CCIATGIAAA CACAATGGG AGCTGAAGGC TIATGGTGCA GGGCTGCTGT CTTCCTACGG	720
10	AGAGCICCIG CACICCCIGI CAGAGGAGCC TGAGGICCCA GCCITTIGACC CAGACACAGC	780
	AGCIGIGCAG COCTACCAAG ATCAAACCTA CCAGCCIGIG TACITIGIGI COCAGAGCIT	840
	CANTGAGGC ANGGACANGC TCAGGANCIA TGCCICIOGI ATCCAGOGCC CATTCICIGI	900
15	GAAGITICAC COGIACACAC ICCOCATICA CGIACICCAC ACCOCICACA COATOCAGO	960
	CICCITEGAG GEGGICCAGG ATGAGCIGCA CACCITEGCC CACGCACTGA GIGCCATTIAG	1020
20	CIAAATGCAT AGGATCC	1037
	(2) INFORMATION FOR SEQ ID NO:18:	
25	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 3425 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: single  (D) TOPOLOGY: linear	
30	(ii) MOLECULE TYPE: UNA (genomic)	
	(iii) HYPOIHETICAL: NO	
35	(iv) ANTI-SENSE: NO	
	(Vii) IMEDIATE SOURCE: (B) CLONE: rTH-IRES-bDBH	
40	(ix) FEATURE:  (A) NAME/KEY: 5'UIR  (B) LOCATION: 16	
45	(ix) FEATURE:  (A) NAME/KEY: excn  (B) LOCATION: 71017	

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	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 10181617	
5	(ix) FEATURE: (A) NAME/KEY: exon (B) LOCATION: 16183411	
10	(ix) FEATURE: (A) NAME/KEY: 3'UIR (B) LOCATION: 34123425	
15	(ix) FEATURE:  (A) NAME/KEY: misc_feature  (B) LOCATION: 10251617  (D) OTHER INFORMATION: /product= "IRES sequence"	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	
	AAGCTTATIGG TOOCCTEGIT COCAAGAAAA GIGIOGGAAT TGGACAAGIG TCACCACCTG	60
	GICACCAAGI TICACCCICA TCIGGACCIG GACCACCCGG GCITICICICA CCAGGIGIAT	120
25	CECCAGCGIC GEAAGCIGAT TGCAGAGATT GCCTTCCAGT ACAAGCACGG TGAACCAATT	180
	COCCATGIGG ANTACACAGC GGAAGAGATT GCTACCIGGA AGGAGGIATA TGTCACCCIG	240
30	AAGEGOCICT ATGCTACOCA TGCCTGCCGG GAGCACCTGG AGEGTTTCCA GCTTCTGGAA	300
	CEGIACIGIG CCIACOGAGA CGACAGCATC CCACAGCIGG AGGACGIGIC CCCCTICTIG	360
	ANGENGUEA CIGGETICCA GEIGGEACCE GIGGEOGGIC TACIGICOGE COGIGATTIT	420
35	CIGGOCAGIC TGGCCITCOG CGIGITICAA TGCACCCAGI ATATCOGCCA TGCCTCCICA	480
	CCTATECATT CACCIGAGOC GEACTECTEC CATGAGCTGT TEGGACATGT ACCCATGTTG	540
40	CCICACOCCA CATTICOCCA GITCICCCAG CACATICGAC TICCATCICT CGGGGCCICA	600
	CATCAACAAA TICAAAAACT CICCACCEIG TACICGITICA CIGICGAATT CCCCIAIGT	66
45	ANACAGNATIG GEGRECIGNA GECTTATIGGT GCAGGGCTGC TGTCTTCCTA CGGAGAGCTC	72
	CIGCACIOCC IGICAGAGGA GOCIGAGGIC CGAGCCITIG ACCCAGACAC AGCAGCIGIG	78
	CAGCCCIACC AAGAICAAAC CIACCAGCCT GIGIACTITIG TGICCGAGAG CITCAATGAC	84

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	GCCAAGGACA AGCICAGGAA CIAIGCCICT CGIAICCAGC GCCCAITCIC TGIGAAGITT	900
5	CACCOGIACA CACIGGOCAT TGACGIACIG GACAGOCCIC ACACCATOCA GOGCIOCITIG	960
	GAGGGGGICC AGGAIGAGCT GCACACCCIG GCCCACGCAC TGAGTGCCAT TAGCTAAATG	1020
	CATAGGATOC GOOGGICTOC CTOCOCCCC CCTAACGITTA CTGGCCGAAG CCCCTTGGAA	1080
	TAAGGOOGET GIGOGITTGT CIATAIGITA TITTOCACCA TATTGOOGIC TITTGGCAAT	1140
	GICAGGOCC GCAAACCIGG CCCIGICITC TICACGAGCA TICCIAGGGG TCITICCCCT	1200
15	CICCOCAAAG GAATGCAAGG TCIGITGAAT GICGIGAAGG AAGCAGITICC TCICGAAGCT	1260
	TOTIGNAGAC ANACANOGIC TGIAGOGACC CITTIGCAGGC AGOGGAACCC COCACCIGGC	1320
	GACAGGIGOC TCIGOGGOCA AAAGOCAOGI GIATAAGATA CACCIGCAAA GGOGGCACAA	1380
20	COCCAGTGOC ACGITIGIGAG TIGGATAGIT GIGGAAAGAG TCAAATGGCT CTCCTCAAGC	1440
	GIATTOWCA AGGGCIGWA GGAIGCOCAG AAGGIACCCC ATTGIATGGG ATCTGATCTG	1500
<b>4</b> 5	GEGCCICGET GCACATECIT TACATGIGIT TAGICGAGGT TAAAAAAACGT CTAGGCCCC	1560
25	CENACONCES CENCEIGEIT TICCITICAN ANACACCATG ATANGCITICC CACANCCATG	1620
	TACGERACIE CEGIEGOCET CITICCIGGIC ATCCICGIGE CIGRACITERA GESCICGECT	1680
30	COORDINATE CONTROL CON	1740
	TOCIGGAACA TOAGCIATICO GOAGGAGACO ATCIACITOC AGCIOCTIGGI GOGGGAGCIC	1800
35	AAGGCIGGIG TOCIGITIGG GAIGIOGGAC OGAGGGGAGC TGGAGAATGC TGACTIGGIG	1860
	GICCICIGEA CIGACAGGEA CGCCCCIAC TITIGGGEATG CCIGGAGIGA CCAGAAGGG	1920
40	CAGGIOCACC TOGACIOCCA GCAGGATIAC CAGCITICIGC GGGCACAGAG GACTOCAGAA	1980
	GEOCIGIACO TECTOTICAA GAGECOTTIT GECACCIGIG ACCOCAAGGA CIACCICATO	2040
	CACCACCO COSTOCACCI GEIGIATICA TICCICEACE ACCOCCICCACE GIOCCICEACE	2100
45	TOTATICAACA CATOOTECTT GOACAGEEG CIGCAGAGEG TGCAGCTGCT GAAGOOCAGC	2160
	ATCCCCAAGC CECCCIECC CECCEACACE CECACCATEE AGATCCCCC CCCCGACGTC	2220

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	CICATOCCOG GOCAGCAGAC CACGIACTICG TECHACGICA COGAGCICCC GGACGGCTIC	2280
	CCCCCCCCACC ACATOGICAT GIACCACCC ATOGICACCG AGGCCAACGA GCCCCTGGTG	2340
5	CACCACATGE AGGICTICCA GIGGGCCCC CAGITICGAGA CCATCCCCCA CITICAGCGCG	2400
	COCTECGACT COAAGATGAA GOOGCAGOGG CTCAACTTCT GOOGTCACGT GCTGGOOGCC	2460
	TEGEOCCIEG GOCCAAGEC CITITIACIAC COAGAGGAAG CAGGOCTEGC CITOCGGGG	2520
10	COORDICTOCT COAGATTTICT COROCTIGGAA GITICACTACC ACAACCCACT GGTGATAACA	2580
	GEOOGGOGG ACTOCTOGGG CATOCGGCTG TACTACACGG CTGCGCTGCG GCCCTTCGAC	2640
15	COCCECATOR TECRECTECS COTECCIAN ACCOUNTINA TECCATOR COCCACEAG	2700
	ACCECTICE TOCICACCES CIACTECACE CACAAGTECA COCACCTECC CCTCOCCCC.	.2760
2.0	TCAGGEATTC ACATCTTOGC CTCTCAGCTC CACACGCACC TGACCGGGGG GAAGGTGGTC	2820
20	ACAGIGCIGG CCAGGGACGG CCGGGACACA GACAICGIGA ACAGGGACÁA CCACIACAGC	2880
	COACACITICC AGGAGATICGS CATGITICAAG AAGGTCGTGT CTGTCCAGCC GGGAGACGTG	2940
25	CICATCACCI CITECACATA CAACACEGAA GACAEGAGGC TEGECCACCGT GEGEGECTIC	3000
	GEGATOCTOG AGGAGATGTG OGTCAACTAT GTGCACTACT ACCOCCAGAC GCAGCTGGAG	306
	CICIGCAAGA GOGOGIGGA COCIGGCIIC CIGCACAAGI ACIICOGOCI CGIGAACAG	3120
30	TICAACAGOG AGGAAGICIG CACCIGOOOC CAGGOGICIG TOOCIGAGCA GITTGOCTOC	3180
	GIECCCIGGA ACICCITICAA COGOGAGGIG CICAAGGCCC IGIAGGCCIT CGCACCCAIC	324
35	TOTATECACT GCAACAGGIC CICCGCCCICCAGG GCGAGIGGAA TOGGCAGCCC	330
	CIECCIGAGA TOGIGIOCAG GITIGGAAGAG COCACOCCIC ACTGOCCAGO CAGOCAGGOT	336
40	CAGAGOCCOG COGGOCCCAC OGTGCTGAAC ATCAGTGGGG GCAAAGGCTG AACGTGGGGG	342
40	GCCCC .	342

## (2) INFORMATION FOR SEQ ID NO:19:

(i) SEQUENCE CHARACTERISTICS:

45

(A) LENGIH: 3432 base pairs (B) TYPE: nucleic acid

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		(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: INA (genomic)	
5	(iii)	HYPOIHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
10	(vii)	IMMEDIATE SOURCE: (B) CLONE: rTHDKS-IRES-bDBH	
15	(ix)	FEATURE: (A) NAME/KEY: 5'UIR (B) LOCATION: 113	
20	(ix)	FEATURE: (A) NAME/KEY: excn (B) LOCATION: 141024	
	(ix)	FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 10251624	
25	(ix)	FEATURE: (A) NAME/KEY: exon (B) LOCATION: 16253418	
30	(ix)	FEATURE: (A) NAME/KEY: 3'UIR (B) LOCATION: 34193432	
35	(ix)	FEATURE:  (A) NAME/KEY: misc_feature  (B) LOCATION: 10321624  (D) OTHER INFORMATION: /product= "IRES sequence"	
40	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:19:	
	AAGCTTO	ECC ACCATEGICC CCIEGITICCC ANGANAGIG TORGANTIGG ACAAGIGICA	60
ΛĒ	COACCIG	FIC ACCAMENTIG ACCCIGATOR GEACCIGAC CACCOGGGOT TOTOTGACCA	120
45	GGIGIAI	CAGOGICOGA AGCIGATIGO AGAGATIGOO TICCAGIACA AGCACGGIGA	180

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	ACCAMPICOC CATGIGGAAT ACACAGOGGA AGAGATIGCT ACCIGGAAGG AGGIATATGT	240
	CACECICAAG GEOCICIAIG CIACOCATEC CIGOOGGAG CACCIGGAGG GITTOCAGCT	300
5	TCTGGAAGGG TACTGTGGCT ACCCAGGAGGA CAGCATOCCA CAGCTGGAGG AGGTGTCCG	360
	CITCITICANG CACCICACCIG CCTTCCACCIT COCACCIG COCACCIC COCACCIG COCACCIO COCACCIC COCACIC COCACCIC COCACCIC COCACCIC COCACCIC COCACCIC COCACCIC COCA	420
	TEATITITICIG GOCAGICIEG OCTIOOGOGI GITICAATEC ACCAGIATA TOOGOCATEC	480
10	CICCICACCT ATGCATTICAC CIGAGOOGA CIGCIGOCAT GAGCIGITIGG GACATGIACC	540
	CATGITIGECT GACGCACAT TIGCCCAGIT CICCCAGGAC ATTIGGACTIG CATCICTGGG	600
15	GECCICAGAT GAAGAAATIG AAAAACICIC CACGGIGIAC TGGITCACIG TGGAATICGG	660
	CCIAIGIAAA CAGAAIGGGG AGCIGAAGGC TIAIGGIGCA GGGCIGCIGI CITCCIAGG	: · 720
	AGAGCICCIG CACICCCIGI CAGAGGAGCC TGAGGICCCA GCCTTTGACC CAGACACAGC	780
20	ACCIGICAG COCTACCAAG ATCAAACCIA CCAGCCIGIG TACTTIGIGT COGAGACCIT	840
	CAATGACGCC AAGGACAAGC TCAAGGAACIA TGCCICICGT ATCCAGGGCC CATTCTCTGT	900
25	CAAGITTICAC COGTACACAC TOGOCATTICA CGTACTOGAC AGOCCICACA CCATOCAGOG	960
	CICCITEGAG GEGGICCAGG ATGAGCIGCA CACCIGGCC CACGCACTGA GIGCCATTAG	1020
	CIAAATCAT ACCATOOCC CCICTOCCIC CCCCCCCC AACGITACIG CCCCAACCC	1080
30	CITEGAATAA GEOOGGIGIG CGITIGICIA TAIGITATIT TOCACCATAT TEOCGICITT	1140
	TEGCAATGIG AGGGCCCGA AACCIGGCCC TGICTICITG ACGAGCATIC CIAGGGGICT	1200
35	TICCCCICIC GCCAAAGGAA TGCAAGGICT GITGAATGIC GTGAAGGAAG CAGITICCICT	1260
	GEAGGITICT TGAAGACAAA CAAGGICIGT AGGEAGCTT TGCAGGCAGC GEAACCCCC	1320
40	ACCIGGOGAC AGGIGOCICT GOGGOCAAAA GOCACGIGIA TAAGATACAC CIGCAAAGGC	1380
40	GECACAACCC CAGIGOCACG TIGIGAGITG CATAGITGIG CAAAGAGICA AATGECTCIC	1440
	CICAAGOGIA TICAACAAGG GGCIGAAGGA IGOOCAGAAG GIACOOCAIT GIAIGGGAIC	1500
45	TGATCIGGG CCICGGIGCA CAIGCITTAC ATGIGITTAG TOGAGGITAA AAAACGICTA	1560
	GCCCCCCCA ACCACGGGGA CGIGGITTIC CITICAAAAA CACGATGATA AGCITGCCAC	1620

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	AMOCATIGNAC GECACOGOGG TEGEOGRICHT OCTGGTCATC CTOGTGGCTG CACTGCAGGG	1680
	CICEGCICCC COCEAGAGC CCITICOCCIT CCACATOCCC CICEACCCC AGGGGACCCT	1740
5	GEAGCIGICO TOGAACATCA GOTATGOGCA GEAGACCATO TACITOCAGO TOCTGGTGOG	1800
	CEACCICAAG CEIGGIGIOC IGITTIGGGAT GIOGGACOGA CEGGACCIGG AGAATICCIGA	1860
1.0	CITIEGIEGIE CICIEGACIE ACAGGGAGG CECCIACITI GEGGAIGCET GGAGIGACCA	1920
	GAAGGGGCAG GTOCACCIGG ACTOCCAGCA GGATTACCAG CITICIGGGGG CACAGAGGAC	1980
	TOCAGNAGE CIGITACCIEC TCITICANGAG GCCTTTTIEGC ACCIGIGACC CCAACGACIA	2040
15	OCICATOGAG GACGGCACCE TOCACCIGGT GIATGGATIC CIGGAGGAGC CGCTCCGGIC	2100
	CCICCAGICC ATCAACACAT CCCCCTTCCA CACCCCCTC CACACCCTCC ACCTCCTCAA	2160
20	CONCACCATO CONTAGORES CONTIGORES GEACAGGGE ACCATGEAGA TOCOGGGCC	2220
	CEACHICCIC ATCCCCCCC ACCAGACCAC GIACTGGTCC TACGTGACCG AGCTCCCCGA	2280
	CESCITOCCC CESCACCACA TOGICATGIA CEAGCOCATC GICACCEAGG GCAACGAGGC	2340
25	CCICGICCAC CACATGGAGG TCTTCCAGIG CECCECCAG TTCCAGACCA TCCCCACTT	2400
	CACCEGGCC TGCCACICCA ACATICAAGCC GCAGCEGCTC AACTTCTGCC GTCACGTGCT	2460
30	CECCECTEG COCCIGERE COMMERCIT TIMEIMOUM GAGGAMETAG COCTEGENT	2520
	CEEEEEECC EECICCICCA CATTICICCE CCIEGAAGIT CACIACCACA ACCCACIEGI	2580
	CATALCAGGC COGCOGCACT CCTCGGGCCAT CCGCCTGTAC TACACGGCTG CGCTGCGGCG	2640
35	CITICEACECE GECATICATES AGCIGGECCT GECGIACACE COOGREATES COATCOOCC	2700
	COAGRAGACE COCTIONICO TOACOGRATIA CIGCACGEAC AAGIGCACCO AGCIGGOCCI	2760
40	GOOGGETCA GEGATTICACA TETTOGECTE TEAGETOCAE ACGEACTICA COEGOOGAA	2820
	GGIGGICACA GIGCIGGOCA GGGACGCGGGGGGGGGGGGGGACA GGGACAACCA	2880
	CIACAGOCCA CACTICCAGG AGATOOGCAT GITGAAGAAG GICGIGICIG TOCAGOOGG	2940
45	AGACGIGGIC AICACCICIT GCACATACAA CACGGAAGAC AGGAGGCIGG CCACCGIGGG	3000

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	GECCTIONES ATOCTOSAGE AGATGIGOST CAACTATIGIG CACTACTACC COCAGACGCA	3060
	GCIGGAGCIC TGCAAGAGG CCGIGGACCC TGGCTTCCIG CACAAGIACT TCCGCCTCGT	3120
5	GAYCAGGITIC AACAGGGAGG AAGICIGCAC CIGCOCCCAG GOGICIGIOC CIGAGCAGIT	3180
	TECCIOCGIG COCTEGNACT CCITCANCOG CGAGGIGCTC ANGROCCIGT ACGGCTTCGC	3240
10	ACCCATICTOC ATGCACTGCA ACAGGTOCTC GGOOGTOOGC TTOCAGGGOG AGTGGAATOG	3300
10	GCAGOOOCIG OCIGAGATOG TGIOCAGGIT GGAAGAGOOC ACOOCTCACT GOOCAGOCAG	3360
	CCAGGCTCAG AGCCCCGCCG GCCCCACCGT CCTGAACATC AGTGGGGGCCA AAGGCTGAAC	3420
15	GIGGGGGC CC	3432
•	(2) INFORMATION FOR SEQ ID NO:20:	•
20	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: CINA	
	(iii) HYPOIHEITCAL: NO	
30	(iv) ANTI-SENSE: NO	
	(VII) IMMEDIATE SOURCE: (B) CLONE: ohPOMC-IRES-070	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
	AGGGCACAGC GGGCCCCCT CCCTCCCCCC	30
40	(2) INFORMATION FOR SEQ ID NO:21:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid	
45	(C) SIRANDEINESS: single (D) TOPOLOGY: linear	

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	(ii) MOLECULE TYPE: CENA	
	(iii) HYPOIHETTCAL: NO	
5	(iv) ANTI-SENSE: NO	
10	(vii) IMMEDIATE SOURCE: (B) CLONE: oIRES-rTHD-071	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
15	CAACCAGGGG ACCATGGTTG TGGCAAGCTT	.30
15	(2) INFORMATION FOR SEQ ID NO:22:	•
20	(i) SEQUENCE CHARACTERISTICS:  (A) LENGIH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	•
25	(ii) MOLECULE TYPE: CLNA	
25	(iii) HYPOIHETICAL: NO	
	(iv) ANTI-SENSE: NO	
30	(Vii) IMEDIATE SOURCE: (B) CLONE: OIRES-rTHD-072	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
	CTIGOCACAA CCATGGICCC CTGGTTCCCA	30
40	(2) INFORMATION FOR SEQ ID NO:23:	
40	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 4499 base pairs  (B) TYPE: nucleic acid	
45	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLFOULE TYPE: DNA (genomic)	

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	(111) HYPOINETICAL: NO	
5	(iv) ANTI-SENSE: NO	
	(vii) IMMEDIATE SOURCE: (B) CLONE: panc-th-dbh fusion	
10	(ix) FEATURE:  (A) NAME/KEY: 5'UIR  (B) LOCATION: 143	
15	(ix) FEATURE:  (A) NAME/KEY: excn  (B) LOCATION: 4489	
20	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 90168	
25	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 169482	
25	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 4831080	
30	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 10812091	
35	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 20922691	
40	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 26924485	
45	(ix) FEATURE:  (A) NAME/KEY: 3'UIR  (B) LOCATION: 44864499	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23	:

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	GOGGOGGGT CACCCCIACA GIOGAGCIGI CACCGICCIT ACAAIGAAAT GOAGCIGGGI	60
_	TATCITCTIC CICATGGCAG TGGITACAGG TAAGGGGCIC CCAAGICCCA AACITGAGGG	120
5	TOCATAMET CIGIGACAGT GECAMICACT TIGOCTTICT TICIACAGG GIGAMITOGG	180
	CITIOCOCC CITOCOCCIG GAGITICAAGA GGGAGCIGAC TGGOCAGOGA CICOGGGAGG	240
10	CHEATGEOCC CEACGEOCCT COCCATIGACE COCCAGGGGC CAGGGCCAC CTGGAGCACA	300
	COCTECTEST CECCECCAS AACAACSACS AGGSCOCCTA CAGGATGGAG CACTTOCGCT	360
	GEGGRAGOCC GOOCAAGGAC AAGGGCIACG GOGGITICAT GACCIOCGAG AAGAGCCAGA	420
15	CECCCTEGT CACCTGTTC AAAAACCCCA TCATCAACAA CECCTACAAG AAGGGCCAGT	• 480
	GAGGGACAG COGGCCCIC TOCCTOCCC COCCTAACG TTACTGGCCG AAGCCCTTG	540
20	CANTANGECC GEIGIGOGIT TEICIATATE TIATTITICCA CCATATTECC GICITTIGEC	600
	AATGICAGG COOGAAACC TGCCCTGTC TTCTTGACGA GCATTCCTAG GGGTCTTTCC	660
	CCICIOGOCA AAGGAATOCA AGGICIGIIG AATGICGIGA AGGAAGCAGT TOCICIOGAA	720
<b>25</b>	GCTTCTTGAA GACAAACAAC GICIGIAGOG ACCCTTTGCA GGCAGOGGAA COCCCCACCT	780
	GEOGRACAGET GOCTICTICOEG OCARARGOCA OGTIGITATIANG ATRICACCITIC ANAGGOGGCA	840
30	CAACCOCAGT GOCACGITGT CAGITGGATA GITGIGGAAA CAGICAAATG CCICICCICA	900
	AGCGIATTICA ACAAGGGCI GAAGGATGOC CAGAAGGIAC COCATTGIAT GGGATCTGAT	960
	CIGGGGCCIC GGIGCACATG CITTACATGT GITTAGICGA GGITAAAAAA CGICIAGGC	1020
35	CCCCCAACCA CGGGGACGIG GITTICCITT GAAAAACACG ATGATAAGCT TGCCACAACC	1080
	ATGGTCCCCT GGTTCCCAAG AAAAGTGTCG GAATTGGACA AGTGTCACCA CCTGGTCACC	1140
40	AAGITICACC CICATCIGCA CCICCACCAC CCCCCCTICT CICACCACGT GIATCCCAG	1200
	OGIOGGAAGO TGATTIGOAGA GATTIGOCTTO CAGIACAAGO ACGGIGAACO AATTICOCOAT	126
A =	GIGGAATIACA CAGOGGAAGA GATTIGCTACC TIGGAAGGAGG TATATIGTICAC GCTGAAGGGC	1320
45	CICIAIGCIA COCAIGOCIG COGGGAGCAC CIGGAGGGIT TOCAGCITICT GGAACGGIAC	138

	IGIGCIAC GARARACAS CAICCALAS CIGARRAGOS IGICCOSCIT CITCARRAGOS	Tado
	OGFACTOGET TOTAGCTGOG ACCOGTGGC GGTCTACTGT COGCOGTGA TTTTCTGGCC	1500
5	AGICIGGOCT TOUGOGIGIT TOAATIGCACC CAGIATATOC GOCATIGOCIC CTCACCTATG	1560
	CATTCACCIG ACCOCCACIG CICCCATGAG CIGITIGGGAC ATGIACCCAT GITIGGCTGAC	1620
10	CECACATTIG COCAGTICIC CCAGGACATT GGACTIGCAT CICIGGGGC CICAGAIGAA	1680
10	GWATTGAAA AACTCTCCAC GETGTACTGG TTCACTGTGG AATTOGGGCT ATGTAAACAG	1740
	AATGGGGAGC TGAAGGCTTA TGGTGCAGGG CTGCTGTCTT CCTACGGAGA GCTCCTGCAC	1800
15	TOOCIGICAG AGRAGOCIGA GGIOOGAGOC TITIGACOCAG ACACAGCAGO TGIGCAGOOC	1860
	TACCAAGATC AAACCIACCA GCCIGIGIAC TITIGIGICOG AGAGCITICAA TGACGCCAAG	1920
	GACAAGCICA GGAACIAIGC CICIOGIAIC CAGOGOCCAT TCICIGIGAA GITIGACOG	1980
20	TACACACIGG CCATTGACGT ACTGGACAGC CCTCACACCA TOCAGGGCTC CTTGGAGGGG	2040
	GICCAGGAIG AGCIGCACAC CCIGGOCCAC GCACTGAGIG CCATTAGCIA AATGCATAGG	2100
25	ATCOGCOCCT CTCCCTCCCC CCCCCTAAC GTTACTGGCC GAAGCCGCTT GGAATAAGGC	2160
	CEGIGIECET TIGICIATAT GITATTITICC ACCATATIEC CEICTITIES CAAIGICAGE	2220
30	GOOGGAAAC CIGGOOCIGT CTICTIGAGG AGCATIOCIA GGGGICITIC COCICIOGOC	2280
30	AMAGGANICC AMEGICIGIT GANIGIOGIG AMEGANGCAG TICCICIGGA AGCITCITGA	2340
	AGACAAACAA OGICIGIAGO GACOCITTIGO AGBCAGOGGA ACOCOCCACO TIGGOGACAGG	2400
35	TECCTCTECTS GOCAMANGOC ACCTGTATAN CATACACCTG CANAGECESC ACANCOCCAG	2460
	TECCACETTE TEAGITECAT AGITETECAA AGAGTCAAAT GECTCTCCTC AAGCGTATTC	2520
40	AACAAGGGC TGAAGGATGC OCAGAAGGTA COCCATTIGTA TGGGATCTGA TCTGGGGCCT	2580
40	OGGIGCACAT CCTTTACATG TGTTTAGTOG AGGITTAAAAA AOGTCTAGGC COCCGAACC	2640
	ACCCCCACAC CATGLACAC CATGATAACAC CATGLACGC	2700
45	ACCEPTED CONTINUE GIVATURE GIRECIGNAL TECAGGERIC GERICOGUE	276
	CACAGOCCE TOCCCITOCA CATOCCCIG CACCOCAGG CCACCCIGCA CCIGIOCIGG	2820

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	AACATCAGCT AIGCGCAGGA GACCATCIAC TICCAGCICC IGGIGCGGGA GCICAAGGCI	2000
5	GETGICCIGT TIGGGAIGIC GGACCGAGGG GAGCIGGAGA AIGCIGACIT GGIGGIGCIC	2940
	TIGGACTICACA COCACOGOCO CITACTITICOG GATIGOCTICGA GTICACCAGAA COCOCAGAA COCOCAGAA	3000
	CACCIGEACT COCAGCAGEA THACCAGCTT CTGCGGGGCAC AGAGGACTCC AGAAGGCCTG	3060
10	TACCIGCICT TCAAGAGGC TITTIGGCACC TGTGACCCCA ACGACTACCT CATCGAGGAC	3120
	GECACOGICC ACCIGGIGIA TEGATICCIG GAGGAGOCC TOCCGICCCT GGAGICCATC	3180
15	AACACATOOG GCTTGCACAC GGGGCTGCAG AGGGTGCAGC TGCTGAAGCC CAGCATOOCC	3240
13	AAGOOGGOC TGOOGGGA CACGOGCACC ATGGAGATOC GOGOCOCCATC	3300
	COCCECCACE ACPACACETA CTEGTECTAC GTCACCCACC TCCCCCACCG CTTCCCCCCC	. 3360
20	CACCACATOG TCATGTAGGA GOCCATOGTC ACCGAGGGCA ACGAGGGGCT GGTGCACCAC	3420
	ATGGAGGICT TOCAGTGGGC GEGGAGTTIC GAGACCATCC GCCACTTCAG GGGGCCTGC	3480
25	CACTOCAAGA TGAAGOOGCA GOGGCTCAAC TTCTGOOGTC ACGTGCTGGC CGCCTGGGCC	3540
27	CIGGGGGCA AGGCCITITA CIACCAGAG GAAGCAGGC IGGCCIICGG GGGGCCCGC	3600
	TOCIOCAGAT TICIOCOCCT GGAAGIICAC TACCACAACC CACIGGIGAT AACAGGOOG	3660
30	COCCACIOCT COGGCATCOG CCIGIACIAC ACGGCIGOGC TGCGGCGCTT CGACGCGGC	3720
	ATCATOGAGC TOGGCCTOGC GTACACGCCC GTGATGGCCA TCCCCCCCCA GGAGACGCCC	3780
35	TICTICTICA COGGETACIG CACGGACAAG TECACCAGE TEGECCTECC CECCTCAGGG	3840
	ATTICACATICT TOGOCTICTICA GCTOCACAGG CACCIGACOG GCCGGAAGGT GGTCACAGIG	3900
	CIGGOCAGGG ACCECCAGA GACAGAGAIC GIGAACAGGG ACAACCACIA CAGOCACAC	3960
40	TICCAGGAGA TOCGCATGIT GAAGAAGGIC GIGICIGICC AGOOGGGAGA OGTGCTCATC	4020
	ACCICITICA CATACAACAC GGAAGACAGG AGGCIGGCCA COGIGGGGGG CITOGGGAIC	4080
45	CIGGAGGAGA TGIGGGICAA CIAIGIGCAC TACIACCCC AGACGCAGCI GGAGCICIGC	4140
	AAGAGOGOG TGGACCCIGG CITOCIGCAC AAGIACTICC GOCIOGIGAA CAGGITCAAC	4200

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	AGGAGAAG TCIGCACCIG COCCAGGGG TCIGICCCIG AGCAGITIGC CICCGIGCCC	4260
	TEGANCIOCT TONACCEGA GETECTONAS COCCIGIACE COTTOCCACO CATOTOCATO	4320
5	CACTECAACA ESTOCIOSEC OSTOCECTIC CAGGGOGAGT GGAATOGGCA GCOCCIGOCT	4380
	GAGATIOGIGI OCAGGITIGGA AGAGOCCACO OCTCACTGOC CAGOCAGOCA GGCTCAGAGC	4440
10	COORCIGET GAACATCAGT GEGEGCAAAG GCTGAACGTG GECGECCEC	4499
10	(2) INFORMATION FOR SEQ ID NO:24:	
15	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANTEINESS: single  (D) TOPOLOGY: linear	
20	(ii) MOLECULE TYPE: cDNA	•
20	(iii) HYPOIHEITCAL: NO	
	(iv) ANTI-SENSE: NO	
25	(Vii) IMEDIATE SOURCE: (B) CLONE: OIRES-074	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
	ANAGOGGOG COCCICIOCC TOCCOCCCC	30
35	(2) INFORMATION FOR SEQ ID NO:25:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDELNESS: single	
40	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
45	(iii) HYPOIHETICAL: NO	
	(iv) ANTI-SENSE: NO .	

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	(vii)	IMEDIATE SOURCE: (B) CLONE: oZeocin-077	
5	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:25:	
		GI CAGICCIGCI CCICCGCCAC	30
10	(2) INFO	EMATION FOR SEQ ID NO:26:	
15	(i)	SEQUENCE CHARACTERISTICS:  (A) IENGIH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: CINA	
20	(iii)	HYPOIHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
25	(vii)	IMEDIATE SOURCE: (B) CLONE: 0IRES-Zeocin-075	
20	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:26:	
30	GGICAACI"	TG GOCATGGTTG TGGCAAGCTT	30
	(2) INFO	RMATION FOR SEQ ID NO:27:	
35	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEINESS: single  (D) TOPOLOGY: linear	
40	,		
	· (11)	MOLECULE TYPE: CINA	
	(iii)	HYPOTHETICAL: NO	
45	(iv)	ANTI-SENSE: NO	

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#### (vii) IMEDIATE SOURCE:

(B) CLONE: oIRES-Zeocin-076

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

#### CTTGCCACAA CCATGGCCAA GTTGACCAGT

30

(2) INFORMATION FOR SEQ ID NO:28:

10

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGIH: 5540 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- 15
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO

20

- (iv) ANTI-SENSE: NO
- (vii) IMEDIATE SOURCE:
- 25 (B) CLONE: POMCDACIH-IRES-THD-IRES-DEH-IRES-Zeocin
  - (ix) FEATURE:
    - (A) NAME/KEY: 5'UIR
    - (B) LOCATION: 1..118

30

40

- (ix) FEATURE:
  - (A) NAME/KEY: exon
  - (B) LOCATION: 119..164
- 35 (ix) FEATURE:
  - (A) NAME/KEY: intron
  - (B) LOCATION: 165..243
  - (ix) FEATURE:
  - (A) NAME/KEY: exon
    - (B) LOCATION: 244..557
    - (ix) FEATURE:
      - (A) NAME/KEY: intron
- 45 (B) LOCATION: 558..1155
  - (ix) FEATURE:

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	(A) NAME/KEY: exon (B) LOCATION: 11562166	
5	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 21672766	
10	(ix) FEATURE:  (A) NAME/KEY: excn  (B) LOCATION: 27674560	
15	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 45615159	
15	(ix) FEATURE:  (A) NAME/KEY: exon  (B) LOCATION: 51605534	
20	(ix) FEATURE: (A) NAME/KEY: 3'UIR (B) LOCATION: 55355540	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
	AAGCTIGGIA COGAGCIOGG ATCCACIAGT AACGGCCCC AGIGIGCIGG AATTCIGCAG	60
	ATATOCATCA CACTGGGGC CGCGTCACCC CTAGAGTCGA GCTGTGACGG TCCTTACAAT	120
30	GAVATICCACC TIGGITTATICT TOTTOCTICAT GECAGTIGGIT ACAGGTAAGG GECTOCCAAG	180
	TOCCAPACIT GAGGGICCAT APACICIGIG ACAGIGGCAA TCACTITIGOC TTTCTTTCTA	240
35	CAGGGGIGAA TICCGCTTTC CCCGGGGTT CAAGAGGGAG CIGACIGGC	300
	AGGEPCIOGS GEPGGEPGAT GEOCOGRAGE GOCCIGOGRA TGAGGEGGA GEGGOCCAGE	360
40	COCPACTICIA COMPAGNICIO CICCICCOCCO COCPACAMENA CENCENCECE COCTACACEA	420
40	TGGAGCACIT COGCIGGGC AGCOCCCCA AGGACAAGOG CIACGGCGGT TICAIGACCT	48
	COCAGAGAGG COAGAGGCC CIGGIGAGGC TGITICAAAAA CGCCATCATC AAGAACGCCT	54
45	ACAAGAAGGG CEAGIGAGGG CACAGOGGGC COCICIOOCT COCCOCOCOC TAACGITACT	60
	מוער לערות הווערות הווערים היותר הווערים היותר איני איני איני איני איני איני איני אינ	66

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	TIGOGICIT TIGECANIGI CAGGGOOGG AAACCIGGO CIGICTICIT CALAGCAIT	120
_	CCIAGGGC TITICCCCICT CGCCAAAGGA AIGCAAGGIC IGITGAAIGI CGIGAAGGAA	780
5	CAGITICCIC TICAACACAA ACAACGICIG TAGOCACCCT TICCAGGCAG	840
	CEGNACCICC CACCIGGGA CAGGIGCCIC TGCGGCCAAA AGCCACGIGT ATAAGATACA	900
10	CCIECANAGE CEGCACANCO CCAGIECCAC GITGIGAGIT CEATAGITGI CEANAGAGIC	960
	ANATOSCICI OCICANGOST ATTOMOCANG GESCIGANGS ATSCOCCAÇAN GETACOCCAT	1020
15	TGIATGGAT CICATCIGGG GOCTOGGIGC ACATGCTTTA CATGIGTTTA GIOGAGGTTA	1080
+0	ANAMAGICI AGGOCCOCG ANCCACGGGG ACGIGGITIT CCTTIGANAA ACACGATGAT	.1140
	AAGCTIGOCA CAACCAIGGT COCCIGGTIC CCAACAAAAG IGICGGAATT GGACAAGIGT	1200
20	CACCACCIEG TCACCAAGIT TGACCCIGAT CIEGACCIEG ACCACCEGE CTICICIGAC	1260
	CAGGIGIATC GOCAGOGICG GAAGCIGATT GCAGAGATTG CCTTCCAGIA CAAGCAGGT	1320
25	GAYCCAATTC COCATGIGGA ATACACAGOG GAAGAGATTG CTACCIGGAA GGAGGIATAT	1380
	GICACCCIGA AGGGCCICIA TGCIACCCAT GCCIGCOGG AGCACCIGGA GGGITTOCAG	1440
	CITCIGGAAC GGIACIGIGG CIACOGAGAG GACAGCATOC CACAGCIGGA GGACGIGIOC	1500
30	OSCITICITICA AGGAGOGGAC TEGGCITOCAG CTGOGACOG TEGGOGGICT ACTGTOCGCC	1560
	OGIGATITIC TOCCAGICI GEOCITOOGC GIGITICAAT GCACOCAGIA TATOCCOCAT	1620
35	GOCTOCTOAC CTATGCATTIC ACCTGAGOOG GACTGCTGOC ATGAGCTGTT GEGACATGTA	1680
	COCATGITGG CIGACOGCAC ATTIGCCCAG TICICCCAGG ACATIGGACT TGCATCICIG	1740
	GEGECCICAG ATGAAGAAAT TGAAAAACTC TOCAGGGGGT ACTGGTTCAC TGTGGAATTC	1800
40	GEGCIATGIA AACAGAATGG GEAGCTGAAG GCTTATGGTG CAGGGCTGCT GTCTTCCTAC	1860
	GEAGAGCICC TGCACICCCIT GICAGAGGAG CCIGAGGICC GAGCCITTICA CCCAGACACA	1920
45	COASCIGICO AGOCCIACOA AGATOAAACO TACCAGOCIG IGIACITIGI GIOCGAGAGO	1980
	TICAAIGAGG CCAAGGACAA GCICAGGAAC TAIGCCICIC GIAICCAGGG CCCATICICI	2040

- 86 -

	GIGNGITIG ACCOGIACAC ACIGGOCAIT GACGIACIGG ACAGOCICA CACAIOCAG	2100
	OCCIOCITICG ACCCCIOCA CEATEACCIG CACACCCICG COCACCCACT CAGICCCATT	2160
5	AGCIPARIGE ATAGRATOGG COCCICIOCO TOCCOCCOC CIRAGGITAC TGGOCGAAGC	2220
	CECTIGGAAT AAGGOOGIG TECGITIGIC TATATGITAT TITICACCAT AITGOOGICT	2280
	TITIGECAATIG TEAGGECOOG GAAACCIGGC CCIGICITCT TEAGGACAT TCCIAGGGT	2340
10	CITICCCCIC TOCCCAAAGG AATGCAAGGT CIGITGAATG TOGIGAAGGA AGCAGITICCT	2400
	CIGGAAGCIT CITIGAAGACA AACAACGICT GIAGCGACCC TITIGCAGGCA GCGGAACCCC	2460
15	CCACCIGGGG ACAGGIGCCI CIGCGGCCAA AAGCCACGIG TATAAGATAC ACCIGCAAAG	2520
	GOGGCACAAC COCAGIGOCA CGITIGIGAGT TIGGATAGITIG TIGGAAAGAGT CAAATIGGCTC	2580
••	TOCICARGOS TATTICARCAR GEGECIGARG GATGOOCAGA AGGIACOCCA TIGIATEGEA	2640
20	TOTGATICTOG GEOCTOGGIG CACATGCTTT ACATGTGTTT AGTOGAGGTT AAAAAACGTC	2700
	TAGECOCCC CAACCACGG CACGIGGITT TOCTTTCAAA AACACGATGA TAAGCTTGCC	2760
25	ACAPOCATET ACCIOCACE GEIGGOGIC TICCIGGICA TOCTOGIGGO TGCACTGCAG	2820
	GETIGEETC COCCUPAÇÃO COCCTICOCO TICCACATOC COCTGEÃCOC CEÃGGGÃCC	2880
30	CIGGAGCIGI CCIGGAACAT CAGCIATGCG CAGGAGACCA TCIACITOCA GCICCIGGIG	2940
30	CEGGACCICA AGGCICGIGI CCIGITIGGG AIGICGGACC CAGGGGACCI CGAGAAIGCI	3000
	GACTIGGIGG TECTCIGGAC TEACAGGAC GEOGOCIACT TIGGGGATEC CIGGAGIGAC	3060
35	CACAAGGGC AGGICCACCT GGACTCCCAG CAGGATTACC AGCITCTGCG GGCACAGAGG	3120
	ACTOCAÇAAG GOCTGTACCT GCTCTTCAAG AGGCCTTTTG GCACCTGTGA COCCAACGAC	3180
40	TACCICATOG AGGAGGCAC OGTOCACCIG GIGIATIGGAT TOCIGGAGGA GOOGCTOOGG	3240
10	TOGCTGEAGT CCATCAACAC ATCCCGCTTG CACACGGGGC TGCAGAGGGT GCAGCTGCTG	3300
	AAGOOCAGCA TOOCCAAGOO GEOCCIGOOC GOEGACAGOO GCACCATEGA GATCOGOOC	3360
45	CONSTRUCTION TO TO TO TO TO THE TOTAL TO THE TOTAL CONTROL OF THE TOTAL CONTROL OT THE TOTAL CONTROL OF THE TOTAL CONTROL OT THE TOTAL	3420
	CACCECTICC CCCCCCACCA CATCCICATG TACCACCCA TCCTCACCCA CCCCAACCAG	3480

- 87 -

	GEGLIGGIGC ACCACATIGGA GGICTICCAG TIGGGCGCCG AGTTCGACAC CATCULCAC	3540
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25	GEGGECTTOG GEATOCIGGA GEAGATGIGC GICAACTATG TGCACTACTA CCCCAGACG	4200
	CAGCIGGAGC TCIGCAAGAG CGCCGIGGACACTICC TGCACAAGIA CITCCGCCTC	4260
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	AGGIGGGGG COCCOCCICT COCTOCCCC COCCTAACGT TACTIGGCGA AGCCCCTTGG	4620
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	ATGIGAGGC COGAAACCT GECCTGICT TCTTGACGAG CATTCCTAGG GGICTTTCCC	4740
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	CITCITICAG ACAAACAACG TCTGTAGCGA CCCTTTGCAG GCAGCGGAAC CCCCACCTG	4860

- 88 -

	COCPACAGGIG CCICIGOGGC CANANGOCAC GIGIATIANGA TACACCIGCA AAGGCGGCAC	4920
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5	GOGIATTICAA CAAGGGCIG AAGGATGOOC AGAAGGIACC CCATTGIATG GGATCTGATC	5040
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10	COCCAPACCAC GEGGACGIGG TITICCITIG AAAAAACACGA TGATAAGCIT GOCACAACCA	5160
10	TEECCAAGIT CACCAGICCC GITCCCGICC TCACCGCCC CCACGICCCC GCAGCCGICC	5220
	AGITICIDEAC CEACOGRETIC GEGITICICCE GEGACTICGI GEAGGACEAC TICECCEGIG	5280
15	TEGICOGGEA CEACGIGACC CIGITICATICA COGOGGICCA CEACCAGGIG GIGCOGGACA	5340
	ACACCCIGGE CIGGGIGIGG GIGGGGGGC TGGACGAGCT GIACGCCGAG TGGICGGAGG	5400
20	TOGIGIOCAC GAACITOOGG GAOGOCTOOG GGOOGGOCAT GAOCGAGATC GGOGAGCAGC	5460
20	OGIGGGGGG GGAGITOGCC CIGGGGGACC OGGCGGCAA CIGGGIGCAC TIOGIGGGGG	5520
	AGGAGGACTOGAG	5540
25	(2) INFORMATION FOR SEQ ID NO:29:	
	(i) SEQUENCE CHARACTERISTICS:	

(A) IENGIH: 829 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOIHETICAL: NO 35

30

- (iv) ANTI-SENSE: NO
- (vii) IMMEDIATE SOURCE: 40

(B) CLONE: ProAKS

(ix) FEATURE:

(A) NAME/KEY: 5'UIR

(B) LOCATION: 1..16 45

(ix) FEATURE:

- 89 -

(A)	NAME/KEY:	exon
(B)	LOCATION:	17820

(ix) FEATURE:

(A) NAME/KEY: 3'UIR

(B) LOCATION: 821..829

### (xi) SECUENCE DESCRIPTION: SEQ ID NO:29:

10 COCAAGCITIC GOCACCATEG CECEGITICCT GACACITTEC ACTIGECTEC TETTECTCEG 60 120 CCCCEGGCIC CIGGOGACCG TGCCGGCCGA ATGCAGCCAG GATTGCGCGA CGTGCAGCTA 15 COECCTAGIG COCCOGGOG ACATCAACTT CCTGGCTTGC GTAATGGAAT GTGAAGGTAA 180 240 ACIGOCITICIT CICAAAATITT GGGAAACCIG CAAGGAGCIC CIGCAGCIGT CCAAACCAGA CCTICCICAA GATGGCACCA CCACCCICAG AGAAAATAGC AAACCGGAAG AAAGCCATTT . 300 20 CCTAGCCAAA ACCTATGGGG CCTTCATGAA AAGGTATGGA GCCTTCATGA AGAAAATGGA 360 TGACCITTAT COCATGGACC CAGAAGAAGA GGCCAATGGA AGTGAGATCC TCGCCAAGCG 420 25 GIATGEGGC TICATGAGA AGGAIGCAGA GGAGGACGAC TOGCIGGCCA ATTOCICAGA 480 CCICCIAAAA GACCIICIGG AAACAGGGGA CAACCGAGGG CGIAGCCACC ACCAGGAIGG 540 CAGIGATAAT GAGGAAGAG TGAGCAAGAG ATATGGGGGC TTCATGAGAG GCTTAAAGAG 600 30 AAGOOOCCAA CIGGAAGAIG AAGOCAAAGA GCIGCAGAAG CGAIAIGGGG GCI'ICAIGAG 660 AAGAGIAGGI COCCAGAGI GGIGGAIGGA CIACCAGAAA CGGIAIGGAG GITICCIGAA 780 35 GOECTTIGOC GAGGCICIGC CCICCGACGA AGAAGGCGAA AGTIACICCA AAGAAGITCC 829 TCANATGCAA AAAACATACG CACCATTTAT CACATTTTAA CCATCCGG

#### (2) INFORMATION FOR SEQ ID NO: 30:

40

(i) SEQUENCE CHARACTERISTICS:

(A) LENGIH: 598 base pairs

(B) TYPE: nucleic acid

(C) STRANDELNESS: single

45 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

540

598

- 90 -

	(iii) HYPOIHETICAL: NO	
5	(iv) ANTI-SENSE: NO	
	(vii) IMEDIATE SOURCE: (B) CLONE: IRES sequence	
10	(ix) FEATURE:  (A) NAME/KEY: intron  (B) LOCATION: 1598	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
	CAATTOOOC CCICIOCCIC CCCCCCCCC AACGITIACTG GCCCAAGCCG CITICGAATAA	60
	GEOGGIGIG CETTIGICIA TAIGITATIT TOCACCATAT TECCGICITT TECCAATGIG	120
20	AGGROSSA AMOCTOGOCC TGTCTTCTTG ACCACCATTC CTAGGGGTCT TTCCCCTCTC	180
	GOCANAGGAA TGCANGGICT GITGANTGIC GTGANGGANG CAGITICCICT GGANGCITCT	240
25	TGAAGACAAA CAAGGICIGI AGOGAGCCIT TGCAGGCAGC GGAACCCCC ACCIGGCGAC	300
	AGGIGCCICT GOGGCCAAAA GOCACGIGIA TAAGATACAC CIGCAAAGGC GGCACAACC	360
	CAGIGOCACG TIGICAGITG CATAGITGIG CAAACAGICA AATGCCICIC CICAAGCGIA	420
30	TICAACAAGG GECIGAAGGA TGOOCAGAAG GIACOCCATT GIATGGGATC TGATCTGGGG	480

35 ACCACEGGA CGIGGITTIC CITICAAAAA CACGAIGAIA AGCITGCCAC AACCAIGG

			1 No.
Applicants or agents file CTI/29	CIP	PCT	international application No.
reference number			

# INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page 54		
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional abeet	
Name of depositary institution		
American Type Culture Colle	ction	
Address of depositary institution finctuains postal code and country		
12301 Parklawn Drive Rockville, Maryland 20852 United States of America Identification Reference by Der	Cell Line, RINa/ProA/	
07 June 1995 (07.06.95)	CRL 11921	
C. ADDITIONAL INDICATIONS (leave plant if not applicab	ic) This information is continued on an additional about	
C. ADDITIONAL INDICATIONS (Leave should for explicable)  In respect of the designation of the EPO, samples of the deposited microorganisms will be made available until the publication of the mention of the grant of the European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, as provided in Rule 28(3) of the Implementing Regulations under the EPC only by the issue of a sample to an expert nominated by requester (Rule 28(4) EPC).  D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designand States)  EPO  E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not explicable)  The moderations insted below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accessed Number of Deposit")		
This sheet was received with the international application  Authorized officer	This sheet was received by the International Bureau on:  Authorized officer	

**SUBSTITUTE SHEET (RULE 26)** 

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Applicant's or agent's tile		International application No.
reference number CTT /29	CIP PCT	!

### INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism reference on page 54 , line S	4-23			
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional abeet			
Name of depositary institution  American Type Culture Called	ction			
Address on depositary institution (including postal code and country)  12301 Parklawn Drive Rockville, Maryland 20852 United States of America Identification Reference by Depos	Cell Line, RINa/ProA/ sitor: P030/P088			
· · · · · · · · · · · · · · · · · · ·	CRL 11921			
07 June 1995 (07.06.95)				
C. ADDITIONAL INDICATIONS (Icave plank if not applicable)	This information is continued on an additional sheet X			
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.				
D. DESIGNATED STATES FOR WHICH INDICATIONS	RE MADE (if the indications ere not for all designated States)			
Finland				
E. SEPARATE FURNISHING OF INDICATIONS (leave blank	k if not applicable)			
The indications listed below will be submitted to the International Bure Number of Deposit?	au later (specify the general nature of the indications e.g., "Accession			
This sheet was received with the international application  Authorized officer  Authorized Authoriz	This sheet was received by the International Bureau on:			

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Applicant's or agent's file CTI/29	CIP	PCT	International application No.

### INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made nellow relate to the microorganism re on page 54	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution  American Type Culture Coll	lection
Address of depositary institution fincluding postal code and country	···
12301 Parklawn Drive Rockville, Maryland 20852 United States of America Identification Reference by Dep	Cell Line, RINa/ProA/
07 June 1995 (07.06.95)	Accession Number CRL 11921
C. ADDITIONAL INDICATIONS (Icave plank if not applicable	r) This information is continued on an additional sheet
Applicant(s) hereby give notice samples of the above-identified only to experts in accordance w Fourth Schedule to the Patents	culture shall be available
D. DESIGNATED STATES FOR WHICH INDICATION	NS ARE MADE (if the indications are not for all designated States)
Singapore	
E. SEPARATE FURNISHING OF INDICATIONS (leave	blank if not applicable)
The indications listed below will be submitted to the Internstional E Number of Deposit?	Buresu later (specify the general nature of the indications e.g., "Accounter
This sheet was received with the international application  Authorized officer  Modelle J. Jan.	For International Bureau use only  This sheet was received by the International Bureau on:  Authorized officer

#### WE CLAIM:

- 1. A cell stably transformed to produce at least one analgesic compound from each of the groups consisting of endorphins, enkephalins, and catecholamines.
- 2. The cell of claim 1, wherein the endorphin is  $\beta$ -endorphin.
- 3. The cell of claim 1, wherein the enkephalin is met-enkephalin.
- 4. The cell of claim 1, wherein the catecholamine is norepinephrine or epinephrine.
- 5. The cell of any one of claims 1-4 wherein the cell is a RIN cell.
- 6. The cell of any one of claims 1-4 wherein the cell is an AtT-20 cell.
- 7. The cell of any one of claims 1-6 wherein the cell additionally produces a compound selected from the group consisting of galanin, somatostatin, neuropeptide Y, neurotensin, or cholecystokinin.
- 8. A cell transformed with a DNA encoding POMC, a DNA encoding TH, a DNA encoding DBH, and a DNA encoding ProA, each DNA molecule operably linked to an expression control sequence.

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- 9. The cell of claim 8 wherein the cell is transformed with pCEP4-POMC-030, pcDNA3-hproA+KS-091, and pZeo-pCMV-rTHAKS-IRES-bDBH-088.
- 10. The cell of claim 8 wherein the cell is transformed with pCEP4-h POMC-ΔACTH-032, pBS-CMV-proA, and pZeo-pCMV-rTHΔKS-IRES-bDBH-088.
- 11. The cell of claim 8 wherein the cell is transformed with pcDNA3-hPOMCDACTH-IRES-rTHD-IRES-bDBH-IRES-Zeocin-073 and pcDNA3-proA+KS-091.
- 12. A transformed cell producing at least one enkephalin, one endorphin and one catecholamine, wherein the cell is transformed with:
- a first vector containing a DNA encoding POMC operably linked to an expression control sequence,
- a second vector containing a DNA encoding pro-enkephalin A operably linked to an expression control sequence,
- a third vector containing a DNA encoding TH operably linked to an expression control sequence and a DNA encoding dopamine beta hydroxylase operably linked to an expression control sequence.
- 13. A method for treating pain comprising implanting at an implantation site in a patient a therapeutically effective number of the cells of any of claims 1-12.

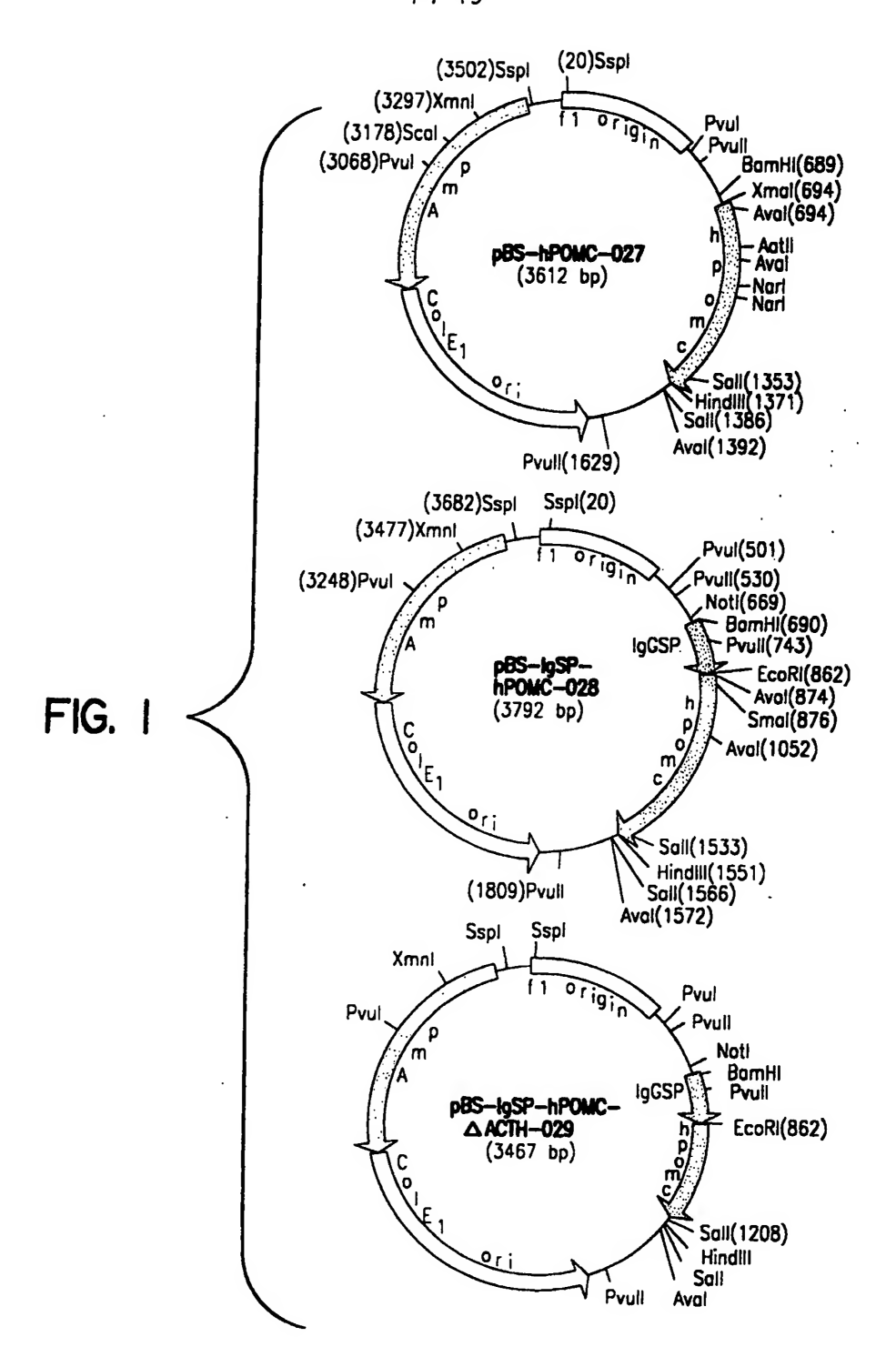
- 14. The method of claim 13 wherein the cells are encapsulated in a semi-permeable membrane to form a bioartificial organ.
- 15. The method of claim 14 wherein the bioartificial organ is immunoisolatory.
- 16. The method of any one of claims 13-15 wherein the implantation site is the CNS.
- 17. The method of any one of claims 13-15 wherein the implantation site is the sub-arachnoid space.
- 18. A method of producing a cell that secretes at least one enkephalin, one endorphin and one catecholamine, comprising transforming the cell with a DNA encoding POMC operably linked to a first expression control sequence, a DNA encoding pro-enkephalin A operably linked to a second expression control sequence, and a DNA encoding TH operably linked to a third expression control sequence and a DNA encoding dopamine beta hydroxylase operably linked to a fourth expression control sequence.
- 19. The method of claim 18 wherein said first, second, third and fourth expression control sequences are identical.

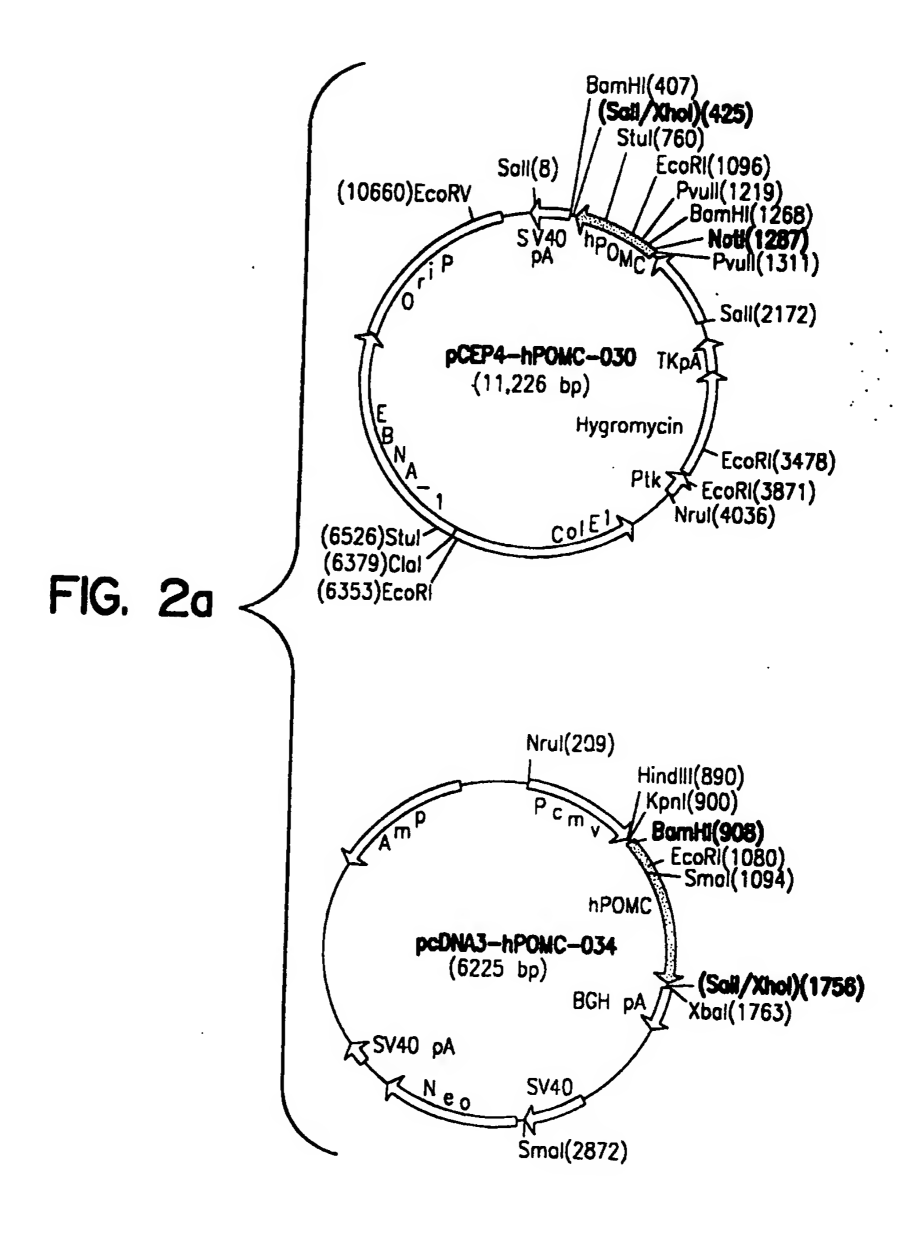
- 94 -

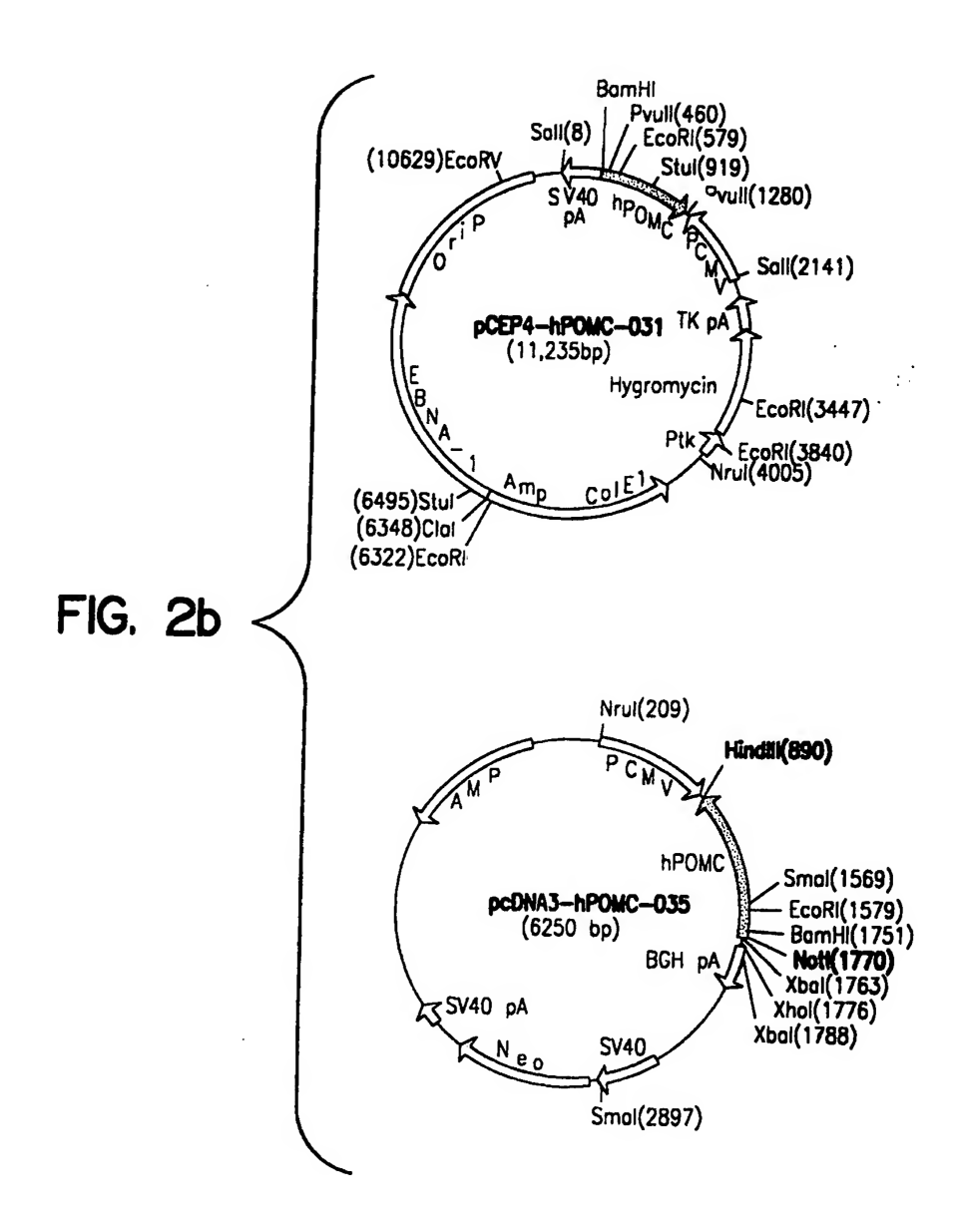
- 20. The use of the cells of any of claims 112 to manufacture a medicant for treatment of pain.
- 21. The cells of claim 20 wherein the cells are implanted.
- 22. The cells of any one of claims 21-22 wherein the cells are encapsulated in a semi-permeable membrane to form a bioartificial organ.
- 23. The cells of claim 22 wherein the bioartificial organ is immunoisolatory.
- 24. The cells of any one of claims 21-23 wherein the implantation site is the CNS.
- 25. The cells of any one of claims 21-23 wherein the implantation site is the sub-arachnoid space.
  - 26. A bioartificial organ comprising:
- (a) a biocompatible, permeable jacket surrounding a core; and
- (b) said core comprising at least one living cell transformed to produce at least one analgesic compound from each of the groups consisting of endorphins, enkephalins, and catecholamines.
- 27. The bioartificial organ of claim 26 for use in treating pain.

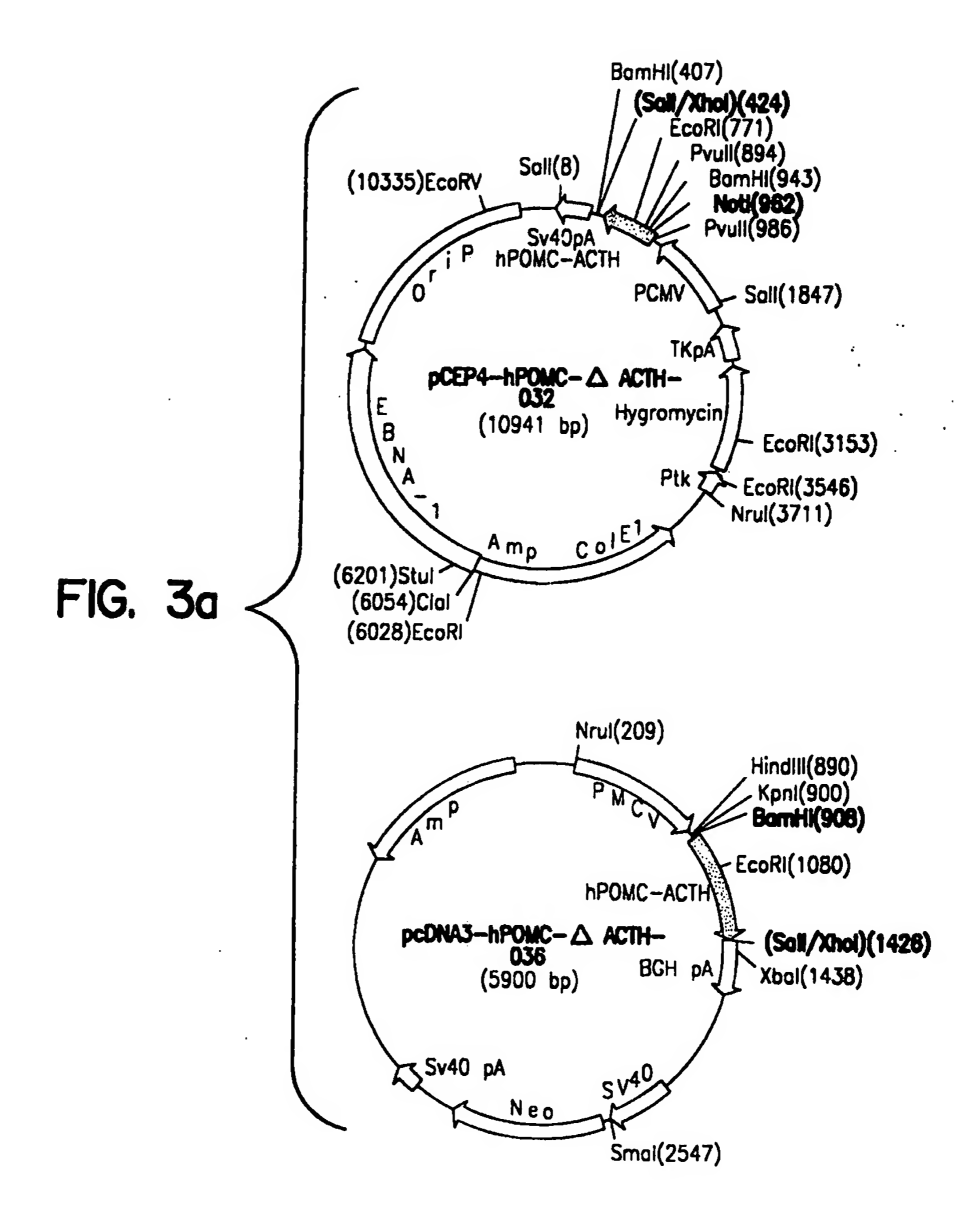
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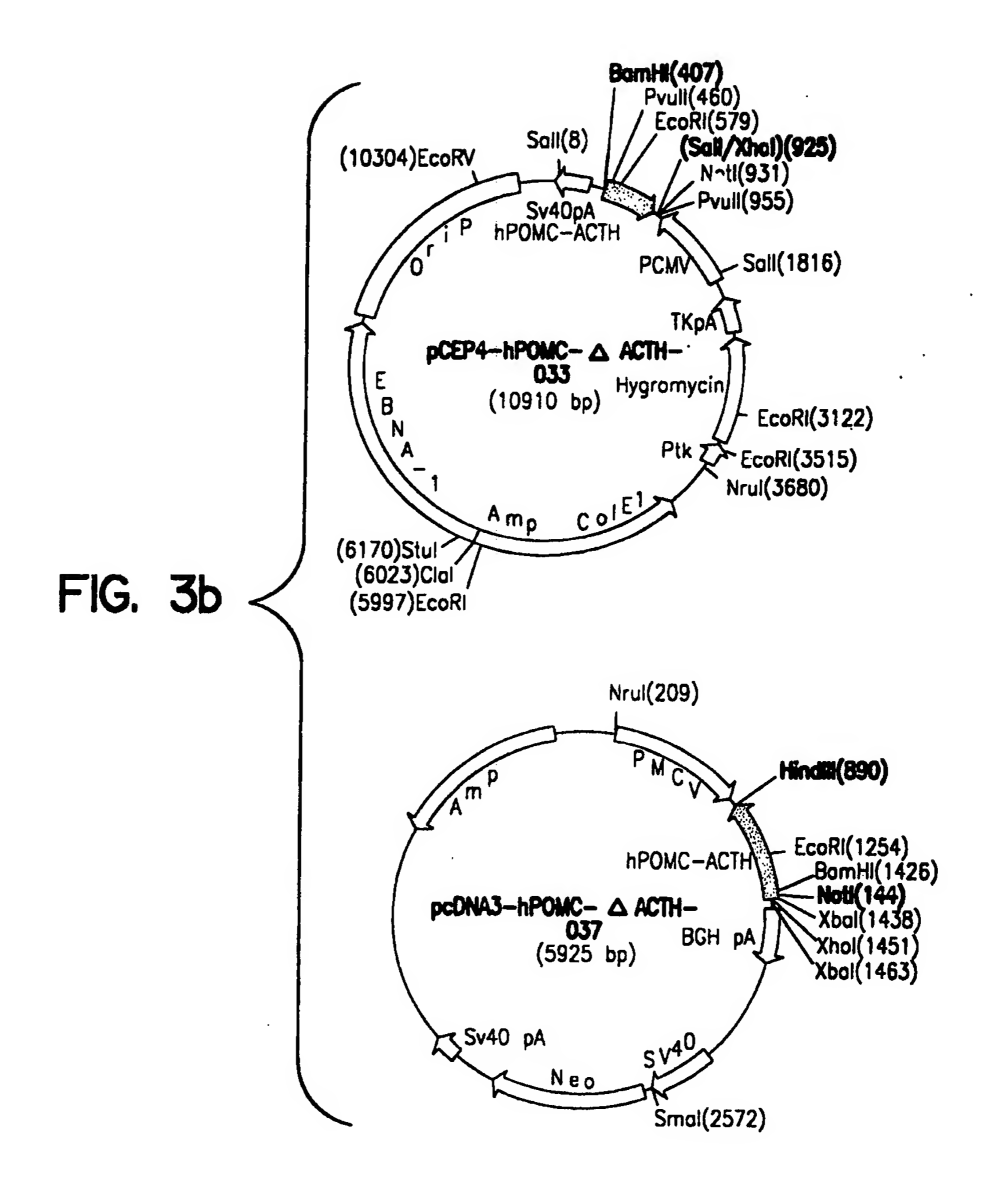
- 28. A method of making a bioartificial organ comprising encapsulating a core comprising at least one living cell transformed to produce at least one analgesic compound from each of the groups consisting of endorphins, enkephalins, and catecholamines, with a biocompatible, permeable jacket.
- 29. The use of a bioartificial organ comprising the cells of claims 1-12 in manufacture of a medicament for treating of pain.

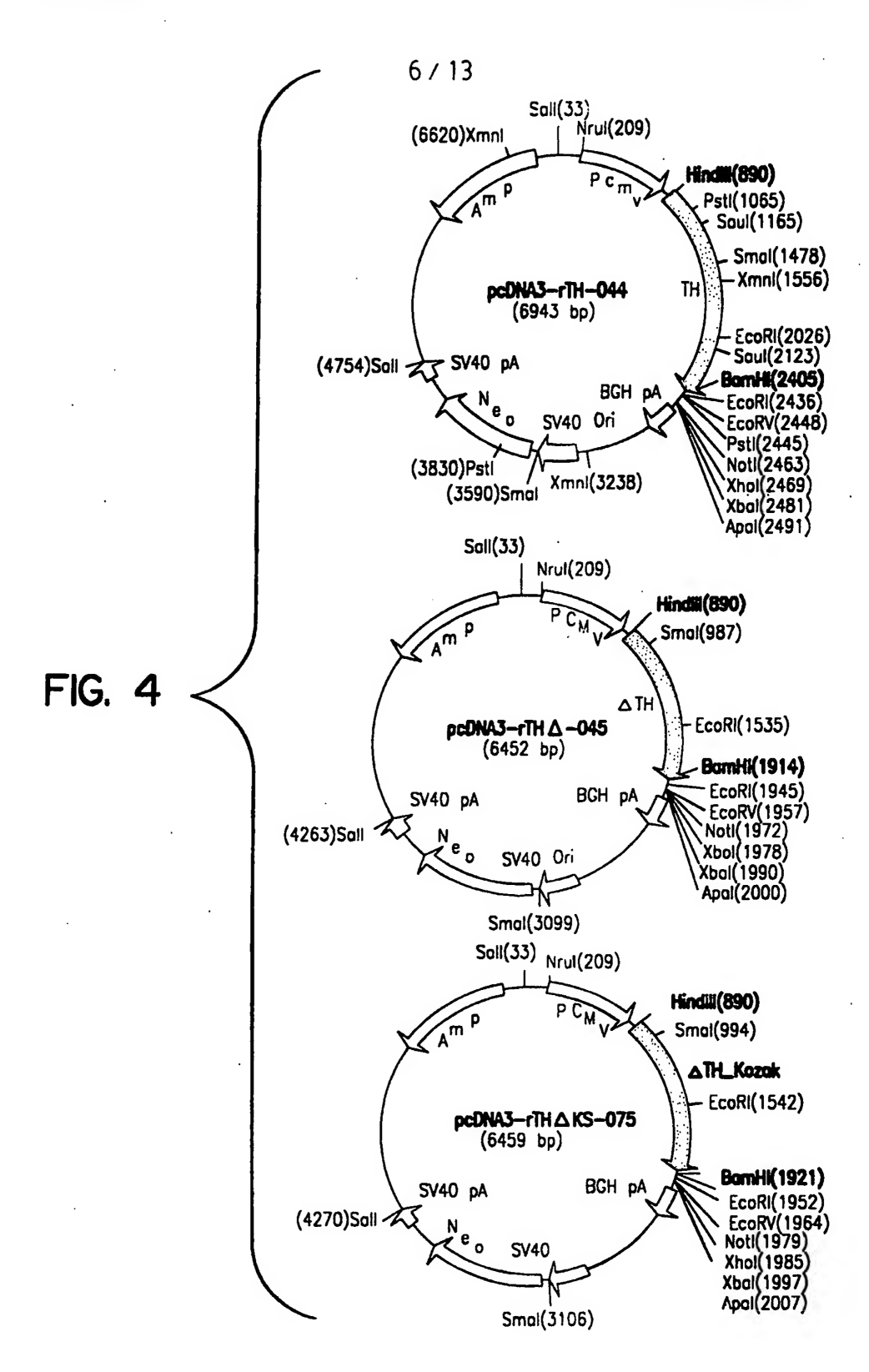












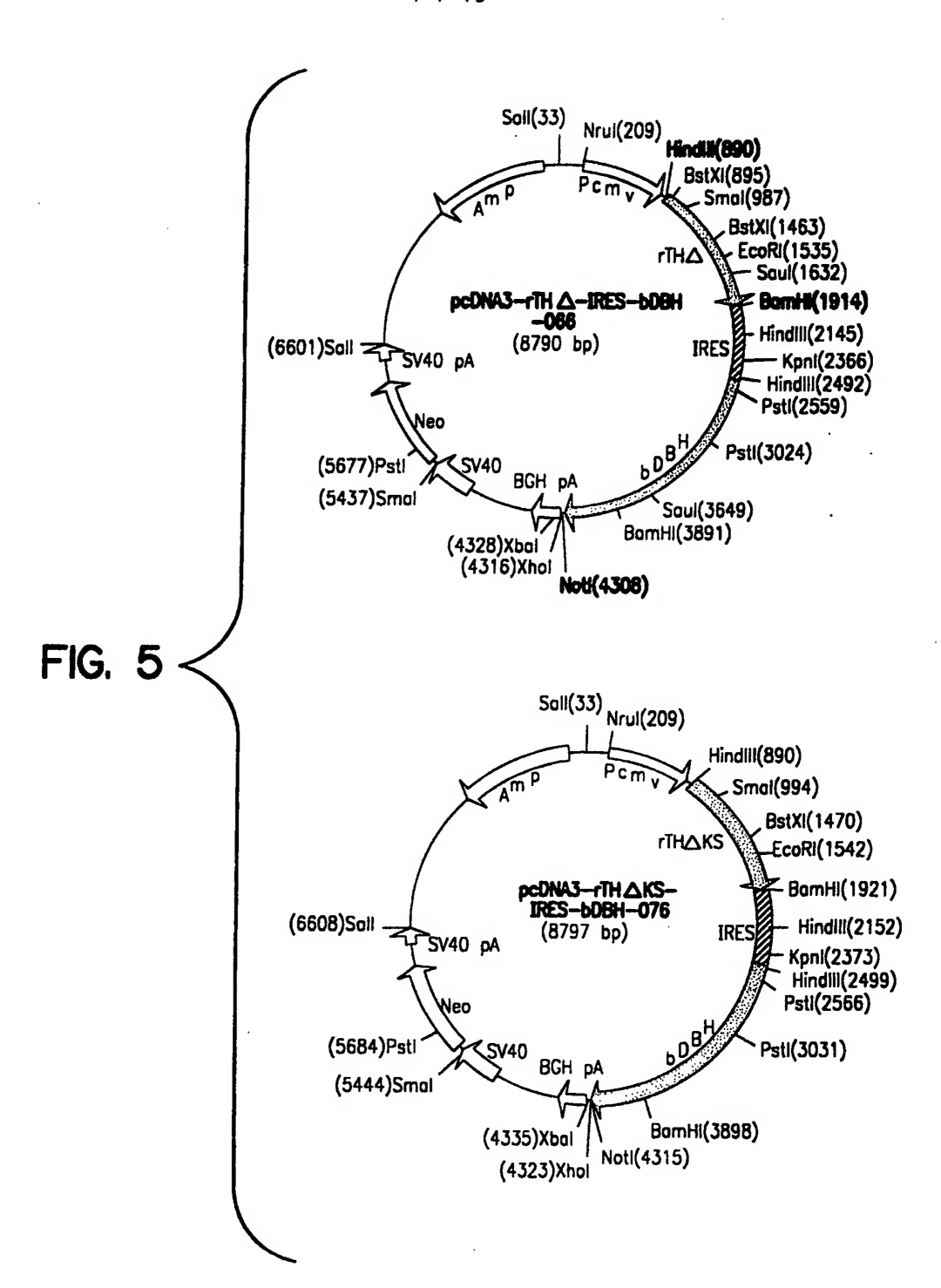


FIG. 6

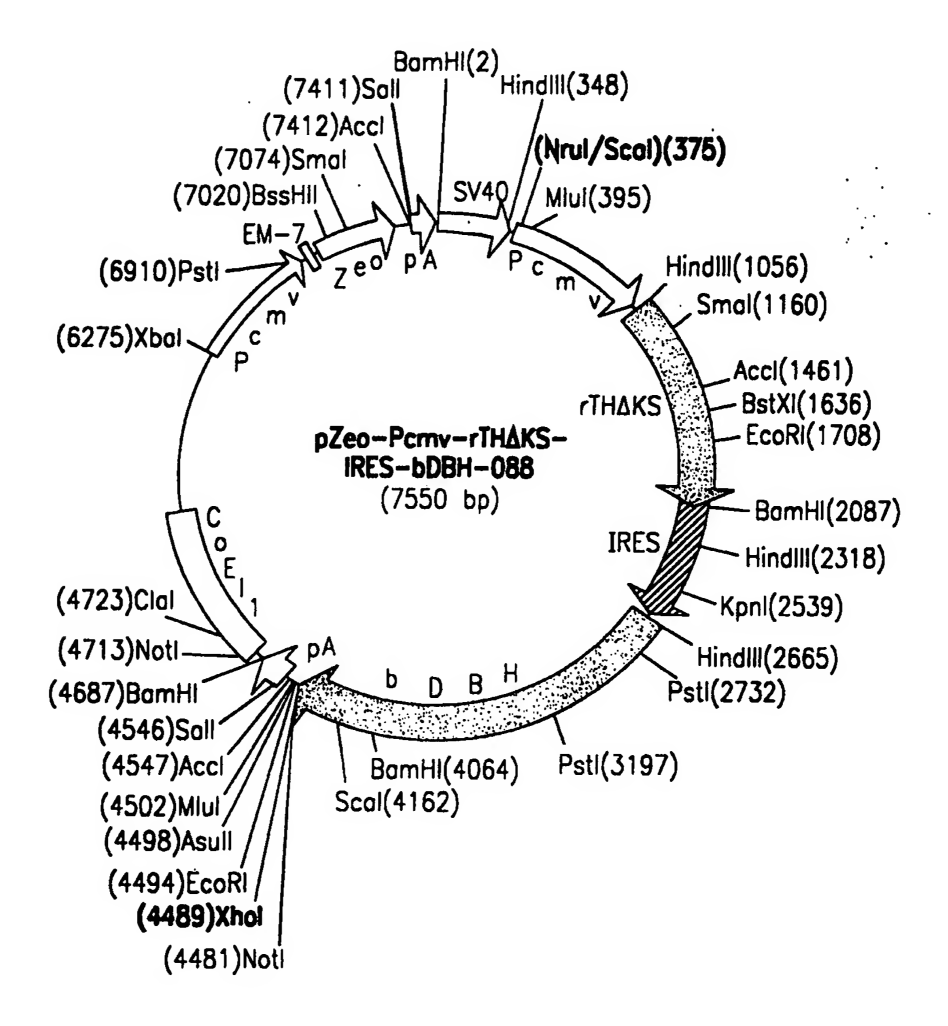


FIG. 7

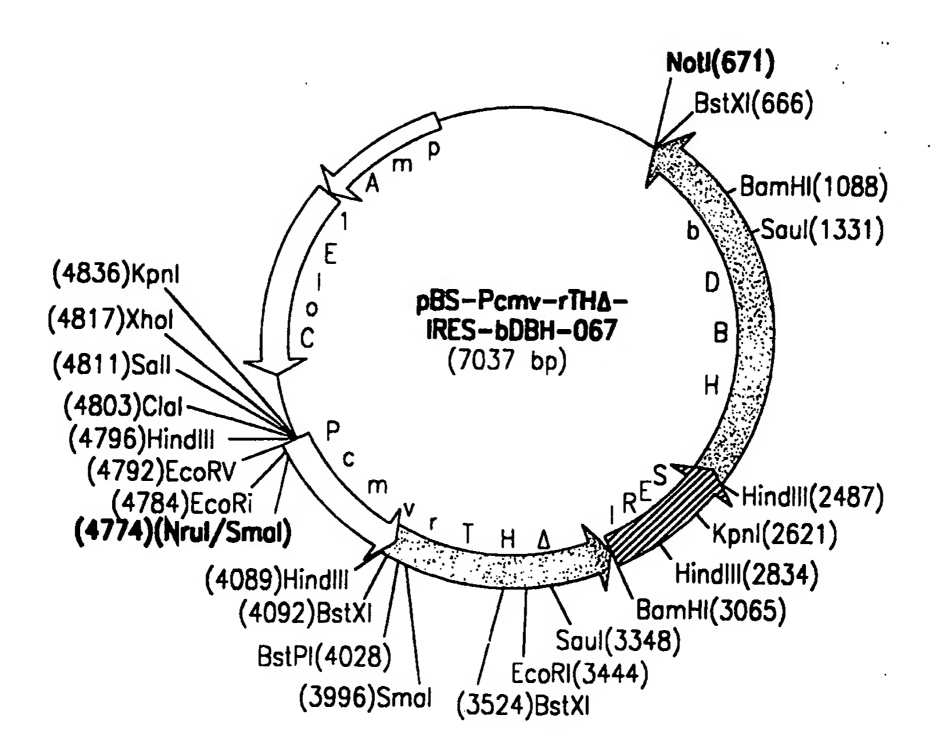


FIG. 8

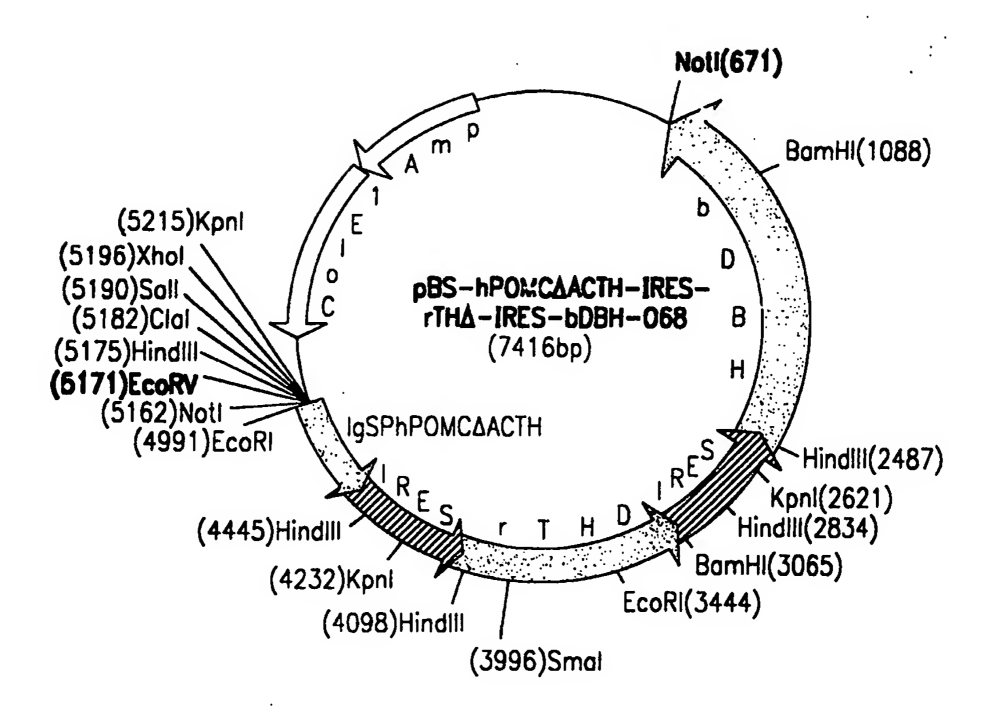


FIG. 9

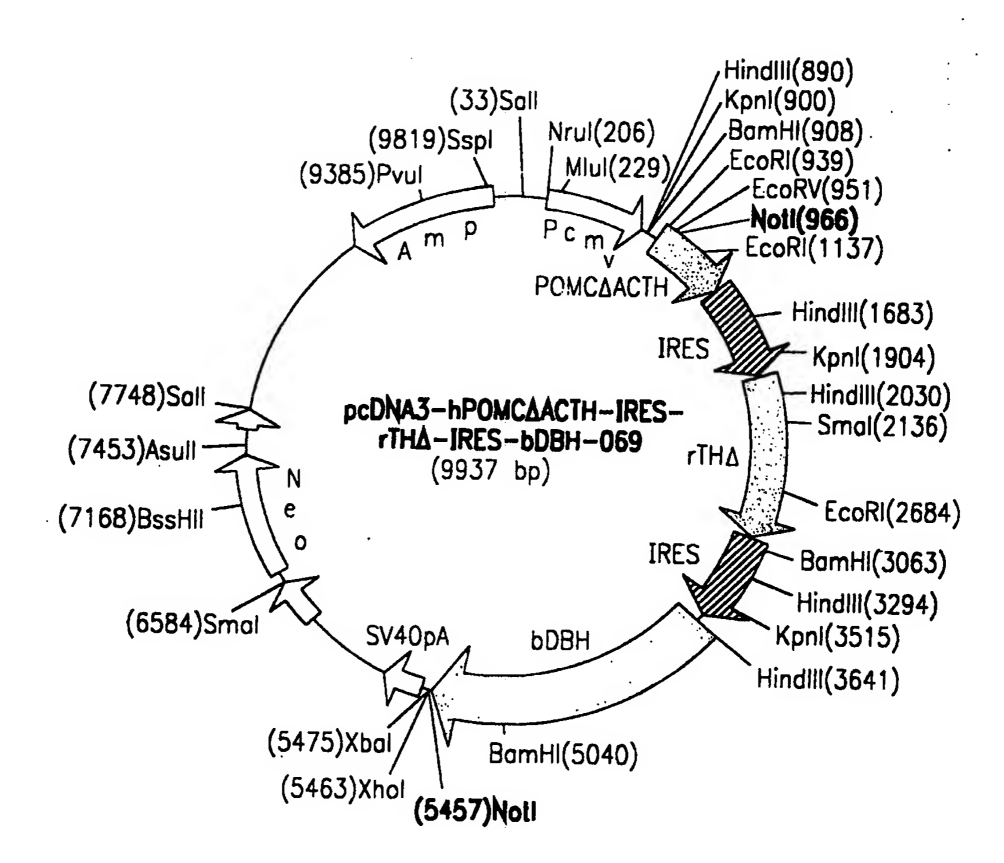
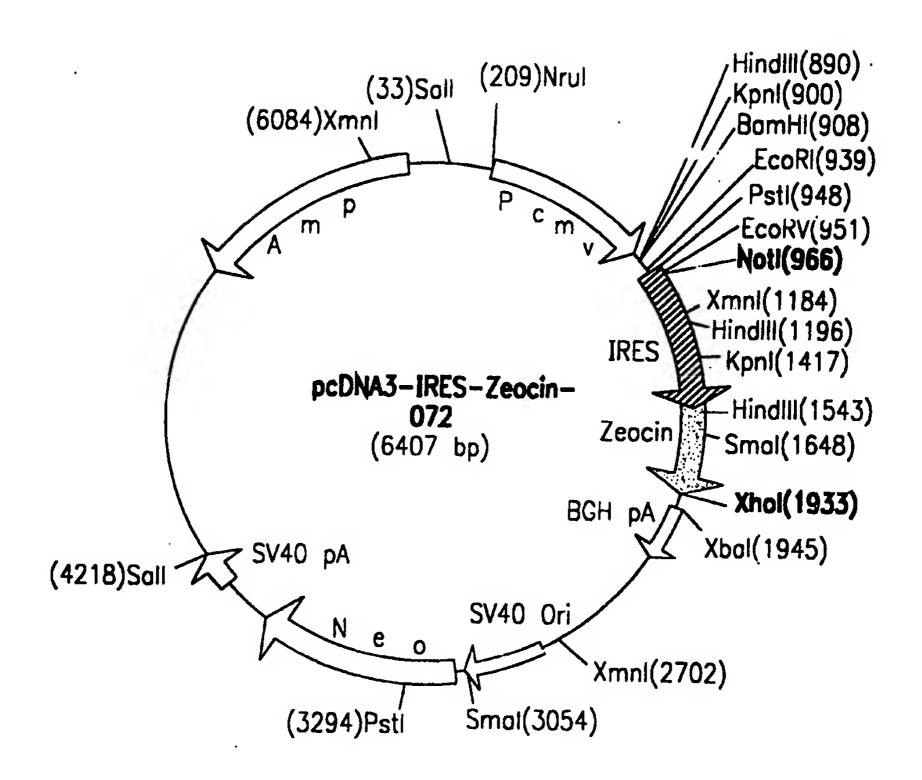


FIG. 10



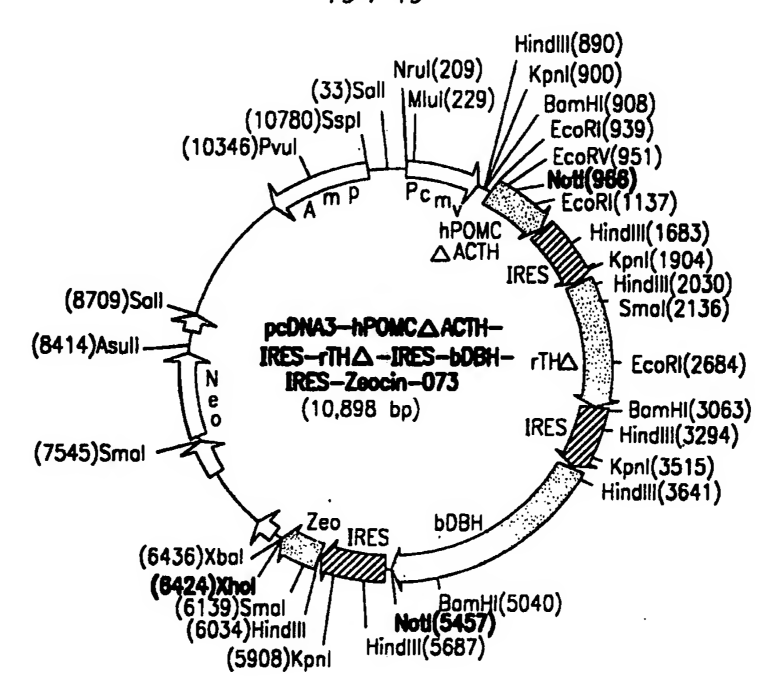


FIG. 11

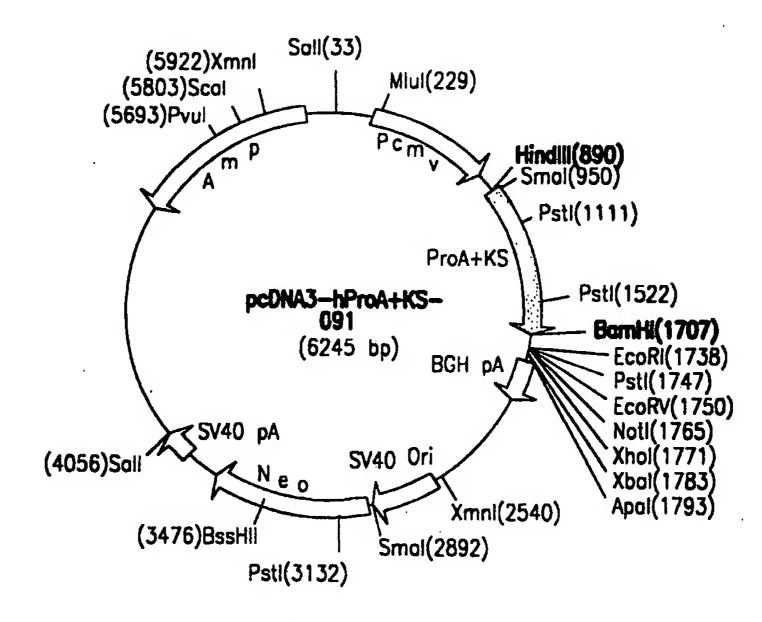


FIG. 12

Is ational Application No PCT/US 96/09629

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	ification of subject matter C12N15/87 C12N5/10	A61K9/48	A61K38/16	A61K38/33
According t	to International Patent Classification (IPC) or to both	national classification	on and IPC	
· · · · · ·	SEARCHED			
	locumentation searched (classification system follows	ed by classification sy	rmbols)	
IPC 6	C12N A61K			
Documenta	tion searched other than minimum documentation to	the extent that such o	locuments are included in	the fields searched
	*** <del>**********************************</del>			
Electronic	tata base consulted during the international search (n	ame of data base and	l, where practical, search t	erms used)
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appro	priate, of the relevan	it passages	Relevant to claim No.
X	WO,A,95 05452 (CYTOTHERAPEUTICS, INC.) 23 February 1995 see the whole document, especially pages 12-31 and Example 6.			1-4,8, 12-29
A	J. NEUROSCI., vol. 14, 1994, pages 4806-4814, XP002018 H.H. WU ET AL.: "Implant or genetically modified A in mouse spinal cord indi antinociception and opioi cited in the application see the discussion.	tation of At AtT-20/hENK uced	cells e"	
[V] Eur	ther documents are listed in the continuation of box (	· [v	Patent family members	and listed in annay
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'A' docum	ntegories of cited documents:  nent defining the general state of the art which is not dered to be of particular relevance		or priority date and not in	fter the international filing date conflict with the application but neighbor or theory underlying the
E carlier	document but published on or after the international date	A 1	document of particular rele cannot be considered nove	evance; the claimed invention
	ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another		involve an inventive step s	when the document is taken alone
citatio	n or other special reason (as specified)		cannot be considered to in	evance; the claimed invention wolve an inventive step when the
	nent referring to an oral disclosure, use, exhibition or means		ments, such combination l	h one or more other such docu- peing obvious to a person skilled
	ent published prior to the international filing date buthan the priority date claimed		in the art. document member of the s	arne patent family
Date of the	actual completion of the international search	1	Date of mailing of the inte	mational search report
1	4 November 1996		2 8.	11. 96
Name and	mailing address of the ISA  European Patent Office, P.B. 5818 Patentiaan 2		Authorized officer	
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,		Yeats, S	
Ĭ.	Fax: ( + 31-70) 340-3016	1		

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_			PC1/US 90/09029		
C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.		
<b>A</b>	PROC. NATL. ACAD. SCI. USA, vol. 83, 1986, pages 7522-7526, XP002018158 J. SAGEN ET AL.: "Analgesia induced by isolated bovine chromaffin cells implanted in rat spinal cord" cited in the application see the abstract and discussion.		1		
A	NATURE, vol. 297, 1982, pages 335-339, XP002018159 M. COCHET ET AL.: "Characterization of the structural gene and putative 5'-regulatory sequences for human proopiomelanocortin" cited in the application see the whole document.		1		
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Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Int	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. <b>X</b>	Claims Nos.: 13-17 because they relate to subject matter not required to be searched by this Authority, namely:  Remark: Although claims 13-17 are directed to a method for treatment of the human body by therapy (Rule 39 PCT), the search has been carried out based on the alleged effects of the composition mentioned in the claims.
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Int	ernational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remari	The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

Information on patent family members

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Patent document cited in search report	Publication date	Patent mem	family ber(s)	Publication date	•
WO-A-9505452	23-02-95	AU-A- CA-A- FI-A- NO-A-	7568094 2169292 960611 960547	14-03-95 23-02-95 09-04-96 12-04-96	
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